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MEG Insight into the Spectral Dynamics Underlying Steady Isometric Muscle Contraction

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To gain fundamental knowledge on how the brain controls motor actions, we studied in detail the interplay between MEG signals from the primary sensorimotor (SM1) cortex and the contraction force of 17 healthy adult humans (7 females, 10 males). SM1 activity was coherent at ~20 Hz with surface electromyogram (as already extensively reported) but also with contraction force. In both cases, the effective coupling was dominant in the efferent direction. Across subjects, the level of ~20 Hz coherence between cortex and periphery positively correlated with the “burstiness” of ~20 Hz SM1 (Pearson r ≈ 0.65) and peripheral fluctuations (r ≈ 0.9). Thus, ~20 Hz coherence between cortex and periphery is tightly linked to the presence of ~20 Hz bursts in SM1 and peripheral activity. However, the very high correlation with peripheral fluctuations suggests that the periphery is the limiting factor. At frequencies <3 Hz, both SM1 signals and ~20 Hz SM1 envelope were coherent with both force and its absolute change rate. The effective coupling dominated in the efferent direction between (1) force and the ~20 Hz SM1 envelope and (2) the absolute change rate of the force and SM1 signals. Together, our data favor the view that ~20 Hz coherence between cortex and periphery during isometric contraction builds on the presence of ~20 Hz SM1 oscillations and needs not rely on feedback from the periphery. They also suggest that effective cortical proprioceptive processing operates at <3 Hz frequencies, even during steady isometric contractions.

Key words: cortex-muscle coherence; corticokinematic coherence; isometric contraction; magnetoencephalography; motor control; primary sensorimotor cortex

Significance Statement

Accurate motor actions are made possible by continuous communication between the cortex and spinal motoneurons, but the neurophysiological basis of this communication is poorly understood. Using MEG recordings in humans maintaining steady isometric muscle contractions, we found evidence that the cortex sends population-level motor commands that tend to structure according to the ~20 Hz sensorimotor rhythm, and that it dynamically adapts these commands based on the <3 Hz fluctuations of proprioceptive feedback. To our knowledge, this is the first report to give a comprehensive account of how the human brain dynamically handles the flow of proprioceptive information and converts it into appropriate motor command to keep the contraction force steady.

Introduction

Steady muscle contraction is maintained by a continuous drive from the cortex to spinal motoneurons and by fine motor adjustments according to proprioceptive feedback (Scott, 2012). Still, we do not know how the brain integrates this proprioceptive feedback to affect motor control.

During steady contraction, the sensorimotor cortical rhythms and the muscle activity, as measured with surface EMG, are cou-
pled at ~20 Hz, a phenomenon known as cortex–muscle coherence (Conway et al., 1995; Salenius et al., 1996, 1997). Some authors suggested that cortex–muscle coherence reflects cortical drive to the muscles (Salenius et al., 1997; Gross et al., 2000), whereas others argued for the existence of a reafferent contribution (Riddle and Baker, 2005; Baker, 2007; Witham et al., 2011) used for sensorimotor control (Baker, 2007). According to this latter view, ~20 Hz sensorimotor oscillations could represent a cortical state that promotes the maintenance of steady motor output (Gilbertson et al., 2005; Androulidakis et al., 2006, 2007; Baker, 2007; Witham et al., 2011). Alternatively, according to a hypothetic mechanism, the sensorimotor system could send pulses at ~20 Hz and monitor the resulting afferent signal to probe the state of the periphery for continuous sensorimotor recalibration (MacKay, 1997; Baker, 2007). Still, these accounts of the functional role of ~20 Hz reafferent signals are highly speculative.

During transient limb movements, cortex–muscle coherence disappears (Kilner et al., 2000) and primary sensorimotor (SM1) activity is coupled with movement kinematics (Kelso et al., 1998; O’Suilleabain et al., 1999; Jerbi et al., 2007; Bourguignon et al., 2011, 2012; Piitulainen et al., 2013a, b). This coupling is known as the corticokinemetic coherence (CKK). When the movement is regular, CKK mainly peaks at movement frequency and its first harmonic (Bourguignon et al., 2011, 2012; Piitulainen et al., 2013a, b). Irregular movements lead to CKK at frequencies <8 Hz (Jerbi et al., 2007). The slow brain activity underpinning CKK is typically strong and reliable enough for brain machine interfaces to use them to detect some features of the movement (Hammon et al., 2008; Waldert et al., 2008; Bradberry et al., 2010; Jerbi et al., 2011). However, CKK appears to reflect the processing of proprioceptive feedback generated by the movements. Indeed, voluntary and passive movements elicit similar CKK level (Piitulainen et al., 2013a), and directionality analyses revealed that the afferent component dominates over the efferent one (Bourguignon et al., 2015).

CKK also unfolds during slow tracking movements. Indeed, the kinematics of slow tracking movements is characterized by fluctuations at frequencies of 1–4 Hz (Craik, 1947), regardless of movement speed (Miall et al., 1986; Roitman et al., 2004; Pasalar et al., 2005), and these fluctuations synchronize with SM1 activity (Dipietro et al., 2011; Hall et al., 2014). In that context, rhythmic fluctuations in movement position are termed submovements (Craik, 1947).

Here, we tested the hypothesis that CKK also manifests itself during isometric steady muscle contractions. To that aim, we analyzed the coupling between MEG signals (Harri and Puce 2017) and natural force fluctuations in an isometric contraction task. First, we expected to uncover significant coupling at ~20 Hz since a previous study revealed the existence of ~20 Hz coherence between MEG and acceleration signals recorded from the index finger in an isometric wrist extension task (Airaksinen et al., 2015). We expected that coupling to dominate in the efferent direction, as is the case for ~20 Hz coupling with EMG. Second, we expected to uncover significant coupling between MEG signals and unavoidable slight fluctuations in contraction force at frequencies <3 Hz (where rhythmic fluctuations in the applied force are the strongest). We expected that coupling to dominate in the afferent direction, as is the case during voluntary movements.

Materials and Methods

Subjects

Seventeen healthy human volunteers (7 females, 10 males; mean ± SD age, 34 ± 7 years; age range, 20–47 years) with no history of neuropsychiatric diseases or movement disorders participated in our study. All subjects were right-handed (mean ± SD score, 90 ± 12; range, 65–100 on the scale from −100 to 100; Edinburgh handedness inventory) (Oldfield, 1971).

The study had a prior approval by the ethics committee of the Aalto University. The subjects gave informed consent before participation, and they were compensated monetarily for travel expenses and lost working hours.

Experimental protocol

Figure 1 illustrates the experimental setup. During MEG recordings, the subjects were sitting with their left hand on the thigh and their right hand on a table in front of them. Subjects’ vision was optically corrected with nonmagnetic goggles (Medigoggles, Cambridge Research Systems) when needed. Subjects were asked to maintain a steady isometric pinch grip of 2–4 N against a custom-made handgrip (connected to a rigid load cell; rigidity 15.4 N/mm; model 1004, Vishay Precision Group) with the right thumb and the index finger (Fig. 1A), and to fixate at a black cross displayed on the center of a screen placed 1 m away in front of them. When the force stepped out of the prescribed limits, a triangle (pointing up or down) appeared on top of the black cross, indicating in which direction to adjust the force, and it disappeared as soon as the force was correctly returned within the limits (Fig. 1B, C). After a ~1 min practice session, two 5 min blocks were recorded, with a minimum of 2 min rest between the blocks. Each block started with ~10 s without contraction, after which subjects were prompted to begin the contraction task. A 5 min task-free block was recorded as well.
Measurements

MEG. The MEG measurements were performed in a three-layer magnetically shielded room (Idemco AG at the MEG Core of Aalto Neuroimaging; http://ani.aalto.fi), Aalto University, with a 306-channel whole-scalp neuromagnetometer (Elekta Neuromag, Elekta Oy). The recording passband was 0.1–330 Hz, and the signals were sampled at 1 kHz. Subjects’ head position inside the MEG helmet was continuously monitored by feeding current into four head-tracking coils located on the scalp: the locations of the coils and at least 200 head-surface points (scalp and nose) with respect to anatomical fiducials were determined with an electromagneto tracker (Fastrak, Polhemus).

EMG and force. Surface EMG was measured from the first dorsal interosseous and the flexor carpi ulnaris of the right hand, which are both recruited during a pinch grip task. EMG electrodes were placed on the muscle bulk, and signals were measured with respect to an electrode placed over the distal radial bone. Recording passband was 10–330 Hz for EMG signals and DC–330 Hz for the force signal. EMG and force signals were then sampled at 1 kHz and recorded time-locked to MEG signals.

MRI. 3D-T1 MRIs were acquired with Signa 3.0 T whole-body MRI scanner (Signa VH/i, General Electric) or with 3T MAGNETOM Skyra whole-body MRI scanner (Siemens Healthcare) at the AMI Centre, Aalto University School of Science.

Data preprocessing

Continuous MEG data were preprocessed off-line with MaxFilter 2.2.10 (Elekta Oy), including head movement compensation. The SSPE preprocessing was applied with a correlated segment length of 0.9 and segment length equal to the recording length (Taulu and Kajola, 2005; Taulu and Simola, 2006). Independent component analysis was then applied to MEG signals filtered through 1–25 Hz, and 1–3 components corresponding to eye-blink and heartbeat artifacts were visually identified based on their topography and time-series. The corresponding components were subsequently subtracted from raw MEG signals.

Functional (nondirectional) coupling

We used coherence analysis to estimate the functional coupling between MEG and all peripheral (2 EMGs, force) signals, and to identify the optimal MEG sensor and muscle for further analyses. Time periods coinciding with visual feedback (i.e., when the force level was not properly kept between 2 and 4 N) were marked as bad to exclude periods during which contraction was intentionally corrected. Time periods for which MEG signals exceeded 5 pT (magnetometers) or 1 pT/cm (gradiometers) were also marked as bad to avoid contamination of the data by any artifact not removed by the preprocessing. Continuous data from the recording blocks were split into 1000 ms epochs with 800 ms epoch overlap (Bortel and Sovka, 2014), leading to a frequency resolution of 1 Hz. Epochs <2 s away from timepoints marked as bad were discarded from further analyses (mean ± SD artifact-free epochs, 2875 ± 130; range, 2573–3031). Coherence spectra were computed between all MEG sensors and nonrectified EMG signals and force signal following the formulation of Halliday et al. (1995), and by using the multitaper approach (5 orthogonal Slepián tapers, yielding a spectral smoothing of ±2.5 Hz) to estimate power- and cross-spectra (Thomson, 1982). Data from gradiometer pairs were combined in the direction of maximum coherence as done by Bourguignon et al. (2015). Then, for each subject and reference signal, the gradiometer pair with the highest coherence value in the 10–30 Hz band was selected among a predefined subset of 9 gradiometer pairs covering the left rolandic region.

A threshold for statistical significance of the coherence (p < 0.05 corrected for multiple comparisons) was obtained as the 95th percentile of the distribution of the maximum coherence (across 10–30 Hz, and across the same 9 gradiometers) evaluated between MEG and Fourier transform surrogate reference signals (1000 repetitions) (Faes et al., 2004). The Fourier transform surrogate of a signal is obtained by computing its Fourier transform, replacing the phase of the Fourier coefficients by random numbers in the range (−π, π), and then computing the inverse Fourier transform (Theiler et al., 1992; Faes et al., 2004). Further analyses were performed for each reference signal with the selected gradiometer signal in the orientation yielding the maximum coherence (MEGSM1). Finally, as EMG was recorded from 2 synergistic muscles, we analyzed only the muscle showing the highest coherence in the 10–30 Hz range (first dorsal interosseous for 10 subjects and flexor carpi ulnaris for the remaining 7 subjects).

Detailed investigations revealed that the ~20 Hz coherence with the force was dampened due to spectral leakage of lower frequencies that were substantially higher in amplitude (see Fig. 1B). For this reason, the coherence was reevaluated based on a procedure less sensitive to spectral leakage as described below. MEG and peripheral signals were filtered through 5-Hz-wide frequency bands centered on 5–40 Hz by steps of 1 Hz. The bandpass filter used in that effect was designed in the frequency domain with zero-phase and 1-Hz-wide squared-sine transitions from 0 to 1 and 1 to 0 (e.g., the filter centered on 20 Hz rose from 0 at 17 Hz to 1 at 18 Hz and ebbed from 1 at 22 Hz to 0 at 23 Hz). Of note, MEG signals were smoothly set to 0 (squared-cosine transition of 1 s) at timings 1 s around artifacts (timepoints of MEG signals exceeding 5 pT for magnetometers or 1 pT/cm for gradiometers); and to avoid edge effects, further analyses were based on timepoints at least 2 s away from artifacts and appearance of visual feedback. The analytical signals were then created by means of the Hilbert transform. From these analytical signals (s(f, t); k = 1, 2 indexing the MEG and the peripheral signal respectively), amplitude spectra were estimated as follows:

\[ A_k(f) = \sqrt{|s(f, t)|^2}, \]

where |·| denotes the absolute value and ⟨·⟩ the average across time. Coherence was then estimated as follows:

\[ \text{Coh}(f) = \frac{\langle |s_1(f, t)\rangle |s_2(f, t)\rangle^2}{\langle |s_1(f, t)|^2 \rangle \langle |s_2(f, t)|^2 \rangle}, \]

where a superscript * denotes the complex conjugate. This procedure consistently enhanced the coherence with the force signal while it left the coherence with EMG signals virtually unchanged.

Finally, we also estimated envelope correlation. To avoid contamination by very slow envelope variations, both envelope signals (|s_k(f, t)|; k = 1, 2) were divided by their low-pass filtered version at 0.1 Hz (squared-sine transition from 0.05 to 0.15 Hz). This procedure is illustrated in Figure 2. Such normalization ensured that the correlation was blind to changes slower than 0.1 Hz that could be linked to, for example, changes in contraction strategy (which induces substantial changes in EMG amplitude). Hence, the envelope coupling was estimated as follows:

\[ R(f)^2 = \text{corr} \left( \frac{|s_1(f, t)|}{|s_1(f, t)|} \right) \cdot \left( \frac{|s_2(f, t)|}{|s_2(f, t)|} \right)^2, \]

where the subscript “lp” stands for “low pass filtered at 0.1 Hz.”

The same analysis was repeated with a finer spectral resolution for the force signal only. In this analysis, amplitude and coherence spectra were computed based on MEG and force signals filtered through 0.6-Hz-wide frequency bands centered on 0.4–40 Hz by steps of 0.2 Hz.

Effective (directional) coupling

We used renormalized partial directed coherence (rPDC) (Schelter et al., 2006) to estimate the causal influence of force and MEGSM1 signals on one another. The estimation of the rPDC requires fitting a multivariate autoregressive model to the data, and the parameters of the model determine the frequency resolution. Here, a multivariate autoregressive model of order 100 was fitted to the data low-pass filtered at 50 Hz and resampled at 100 Hz with the ARfit package (Schneider and Neumaier, 2001). Across subjects and conditions, the optimal model order range was 17–34 (mean ± SD, 21 ± 4) according to Schwarz’s Bayesian criterion and 46–376 (166 ± 97) according to Akaike’s final prediction error, both implemented in the ARfit package (Schneider and Neumaier, 2001). Adopting a model order of 100 therefore represented a good compromise between the two criteria. The chosen parameters (resampling and model order) enabled us to explore frequencies up to 50 Hz with a 1 Hz frequency resolution. Finally, to achieve a frequency smoothing similar to that of coherence spectra, rPDC was smoothed with a square
Figure 2. Illustration of the procedure used to obtain envelopes of band-limited signals corrected for slow drifts. A, Raw MEG signal recorded over the primary sensorimotor cortex (MEGSM1). B, Band-limited MEGSM1 envelope. That is, the Hilbert envelope (black trace) of MEGSM1 signal filtered through a narrow band (gray trace). In the present illustration, the band was 5 Hz wide and centered on 20 Hz (hence the ~20 Hz MEGSM1 envelope). The same procedure was repeated for center frequencies from 5 to 40 Hz by steps of 1 Hz (data not shown on the figure). C, ~0.1 Hz fluctuations of the envelope displayed in B. D, Envelope corrected for slow drifts. That is, the envelope (displayed in B) divided timepoint by timepoint by its slow fluctuations (displayed in C). The coefficient of variation (CoV) of that envelope (CoV-E; computed over 10 min) is indicated in the top right corner. E–H, Same as A–D for the EMG signal. I–L, Same as A–D for the force signal. M–P, Same as A–D for a white noise. All corrected envelopes fluctuate around 1, but qualitatively less so for the white noise, leading to a lower CoV.
kernel wide of 5 frequency bins, following the approach proposed by Sommerlade et al. (2009). The ensuing spectral smoothing was ±2.5 Hz. This procedure yielded for each subject one rPDC spectrum in the efferent direction (MEGSM1 → force) and one in the afferent direction (force → MEGSM1).

**Link between cortex–muscle coherence and burstiness of brain and peripheral signals**

We performed additional analyses to clarify the causal influence of ~20 Hz fluctuations in SM1 and peripheral signals on one another. We know that activities of both the SM1 cortex and periphery (EMG, force) are characterized by bursts of ~20 Hz cycles often followed by ~1-s-long silent periods (Jasper and Penfield, 1949; Murthy and Fetz, 1992, 1996; Baker et al., 1997; Gilbertson et al., 2005). We also know that the ~20 Hz cortex–muscle coherence is tightly linked to the presence of these ~20 Hz bursts in both SM1 and peripheral signals (Mcauley et al., 1997; Kilner et al., 2000, 2003; Gilbertson et al., 2005; Kristeva et al., 2007; Ushiyama et al., 2011; Matsuya et al., 2013). Accordingly, we here quantify the “burstiness” of MEGSM1 and peripheral signals, and strive to ascribe the interindividual variability in this burstiness to that in ~20 Hz coherence between MEGSM1 and peripheral signals. Of note, the magnitude of the cortex–muscle coherence is widely known for its great interindividual variability, from values below the detection limit afforded by even 10-min-long recordings (i.e., ~<0.003) up to values of 0.3 (Pohja et al., 2005; Bayraktaroglu et al., 2013).

Figure 2 illustrates the processing procedure we used (detailed below) to estimate the “burstiness” of MEGSM1 and peripheral signals. The coefficient of variation (CoV; ratio between the SD and the mean) of the envelope of a bursting signal is higher than that of white noise filtered similarly (in our setting, the CoV of a filtered white noise is ~0.51). Accordingly, we filtered MEGSM1 and peripheral signals through 5-Hz-wide frequency bands centered on 3–40 Hz by steps of 1 Hz, and computed their CoV-E. To avoid contamination by very slow amplitude variations, amplitude signals were divided by their low-pass filtered version at 0.1 Hz before estimating the CoV.

A threshold for statistical significance of maximal CoV-E value across 10–30 Hz (p < 0.05 corrected for multiple comparisons, one threshold per subject and per signal: MEGSM1, force, and EMG) was obtained as the 95th percentile of the distribution of the maximum CoV-E, across 10–30 Hz, of white noise signals (1000 repetitions) (Faes et al., 2004).

We searched for an association between (1) the magnitude of ~20 Hz cortex–muscle coherence (maximum across 10–30 Hz) and (2) the CoV of ~20 Hz MEGSM1 and peripheral signals (maximum across 10–30 Hz) with Spearman and Pearson correlation (across the 17 subjects). Pairs of Pearson correlation coefficients were compared with the Steiger test (Steiger, 1980).

We also computed the sensor topography for the CoV-E of ~20 Hz MEG signals. Practically, for each gradiometer pair, we retained the maximum CoV-E of the MEG signals (1) in 20 different directions (from 0 to π) within the 2D space spanned by the two gradiometers, and (2) filtered through 5-Hz-wide frequency bands centered on 18–22 Hz by steps of 1 Hz.

**Coupling at lower frequencies**

Figure 3 presents representative samples of the slow fluctuations of MEGSM1 and force signals considered here.

We estimated the coherence at lower frequencies (<5 Hz) between MEG signals (Fig. 3B) and force fluctuations (Fig. 3E). The analysis was identical to that described in Coherence analysis, except that epochs were 5000 ms long, affording a finer spectral resolution of 0.2 Hz and 3 tapers were used (yielding a spectral smoothing of ±0.3 Hz). In addition, MEG and force signals were high-pass filtered at 0.2 Hz to avoid spectral leakage from near to DC components. Another difference was that the threshold for statistical significance (p < 0.05 corrected for multiple comparisons) was obtained for the mean coherence across 0.5–3 Hz.

![Figure 3](image-url)
The same analysis was repeated to evaluate the coupling between MEG signals and fluctuations in the absolute change rate of the force (Fig. 3F).

The absolute change rate of the force was obtained as the absolute value of the time derivative of the force signal band-passed through 0.5–10 Hz. It is relevant to seek coupling with this signal given that some proprioceptive receptors (e.g., Golgi tendon organs) are sensitive to the force itself, whereas others (e.g., the primary endings of muscle spindles) are sensitive to the change rate of muscle stretch.

The same analysis was repeated also to evaluate the coupling between the envelope of MEG signals filtered through 15–25 Hz (Fig. 3C) and the slow fluctuations of both the force signal and its absolute change rate. In what follows, the envelope of MEG signals filtered through 15–25 Hz will be referred to as the ~20 Hz MEG envelope. This envelope signal fluctuates at frequencies <10 Hz.

We used rPDC to quantify the causal influence of the following: (1) the envelope of ~20 Hz MEGSM1 signals, (2) the MEGSM1 signals as such, and (3) the force on one another. A multivariate autoregressive model of order 50 was fitted to the data low-pass filtered at 5 Hz and resampled at 10 Hz, enabling us to explore frequencies up to 5 Hz with a 0.2 Hz frequency resolution. Across subjects and conditions, the optimal model order range was 2–6 (mean SD, 4 ± 1.5) according to Schwarz’s Bayesian criterion and 59–147 (96 ± 25) according to Akaike’s final prediction error. To achieve a frequency smoothing similar to that of coherence, rPDC was smoothed with a square kernel wide of 3 frequency bins, leading to a spectral smoothing of ±0.3 Hz. This procedure yielded, for each subject, 6 rPDC spectra: one for each possible combination of directions (2 possibilities) and signal pairs (3 possibilities).
The same procedure was repeated to quantify the causal influence of the following: (1) the envelope of $-20\,\text{Hz}\,\text{MEG}_{\text{SM1}}$ signals, (2) the MEG$_{\text{SM1}}$ signals as such, and (3) the absolute change rate of the force on one another.

Finally, we used temporal response functions (TRFs) to model how force signals affect the temporal dynamics of $-20\,\text{Hz}\,\text{MEG}_{\text{SM1}}$ envelope and MEG$_{\text{SM1}}$ signals. A similar approach has been used previously to model brain responses to continuous speech sounds (Lalor and Foxe, 2010; Zion Golumbic et al., 2013; Lawler et al., 2015). TRFs are the direct analog of evoked responses in the context of continuous stimulation.

Practically, we used the mTRF toolbox (Crosse et al., 2016) to estimate the TRF of MEG$_{\text{SM1}}$, and of $-20\,\text{Hz}\,\text{MEG}_{\text{SM1}}$ envelope associated with force signals, all signals being filtered through 0.5–5 Hz and downsampled to 20 Hz. For each subject, the TRFs were modeled from $-1.5$ to $2.5\,\text{s}$, for a fixed set of ridge values ($\lambda = 2^0, 2^1, 2^2, \ldots, 2^{20}$). We adopted the following 10-fold cross-validation procedure to determine the optimal ridge value: For each subject, TRFs were estimated based on 90% of the data. TRFs were then multiplied by a window function with squared-sine transition on the edges (rising from 0 at $-1.5$ to 1 at $1\,\text{s}$ and ebbing from 1 at 2 to 0 at $2.5\,\text{s}$) to dampen regression artifacts. We next used these windowed TRFs to predict the 10% of data left out, and estimated the Pearson correlation coefficient between predicted and measured signals. These correlation values in the 10 runs were tested against 0 with a Wilcoxon signed rank test. Finally, the square of the mean correlation value across the 10 runs provided an estimate of the proportion of variance explained by the signals imputable to force fluctuations. In the final analysis, we used the ridge value maximizing the mean explained variance across our 17 subjects (sum of the logarithm across the 2 signals considered: MEG$_{\text{SM1}}$, and $-20\,\text{Hz}\,\text{MEG}_{\text{SM1}}$ envelope; $\lambda = 2^{15}$).

The same procedure was repeated to estimate the TRFs to the absolute change rate of the force of MEG$_{\text{SM1}}$, and $-20\,\text{Hz}\,\text{MEG}_{\text{SM1}}$ envelope, all signals being filtered through 0.5–5 Hz.

**Impact of heartbeats on low-frequency coupling**

Heartbeats transiently increase blood pressure in all blood vessels of the body, leading to changes in the pressure measured at the fingertips (Parati et al., 1989). Such transient pressure changes in the fingertips are thus expected to slightly impact the contraction force at frequencies matching cardiac rhythm and its harmonics, and they also produce some heartbeats-locked changes in the proprioceptive afference because of matching cardiac rhythm and its harmonics, and they also produce some about coherence at frequencies closest to individual heart rate and its harmonics ($f_{\text{heart-rate}}$) with that at frequencies farthest from heart rate and its harmonics ($f_{\text{off-heart-rate}}$). To that aim, and for each subject, we estimated the minimum distance from each frequency bins in 0.5–3 Hz to heart rate and its harmonics. The first half of the frequency bins of minimum distance were assigned $f_{\text{heart-rate}}$ and the other half to $f_{\text{off-heart-rate}}$. For each subject, we then evaluated the mean coherence at $f_{\text{heart-rate}}$ and $f_{\text{off-heart-rate}}$ separately and compared both measures between subjects with a Wilcoxon test.

**Source reconstruction**

We estimated the cortical distribution of the coherence phenomena highlighted by the sensor-level analyses. To that aim, individual MIRIs were first segmented using Freesurfer software (Martinos Center for Bio-medical Imaging, Charlestown, MA; RRID:SCR_001847) (Reuter et al., 2012). MEG and segmented MRI coordinate systems were coregistered using the three anatomical fiducial points for initial estimation and the head-surface points to manually refine the surface coregistration. Then, the MEG forward model based on a one-shell boundary element model of the intracranial space was computed for three orthogonal current dipoles placed on a homogeneous 5 mm grid source space that covered the whole brain (MNE suite; Martinos Center for Biomedical Imaging, Charlestown, MA; RRID:SCR_005972) (Gramfort et al., 2014). To simultaneously combine data from the planar gradiometer and the magnetometer sensors for source estimation, sensor signals and the corresponding forward-model coefficients were normalized by their root-mean-square noise estimated from the task-free data filtered through 1–195 Hz. For each source, the forward model was then reduced to its two principal components of highest singular value, which closely correspond to sources tangential to the skull. Source time courses were then estimated from the sensor data with a minimum-variance beamformer built with the covariance matrix of the task-free data filtered through 1–195 Hz (Van Veen et al., 1997). Maps of the parameters estimated in the sensor space were estimated similarly in the source space (source pairs taking the role of the gradiometer pairs). Then, a nonlinear transformation from individual MRIs to the standard MNI brain was computed using the spatial-normalization algorithm implemented in Statistical Parametric Mapping (SPM8; Wellcome Department of Cognitive Neurology, London; RRID:SCR_007037) (Ashburner et al., 1997; Ashburner and Friston, 1999) and applied to individual maps. Finally, group-level maps were obtained by averaging normalized maps across subjects.

**Results**

**Coupling with EMG and force at $-20\,\text{Hz}$**

MEG oscillations measured from the contralateral (left) SM1 cortex (MEG$_{\text{SM1}}$) were coherent at $-20\,\text{Hz}$ with the surface EMG (Coh$_{\text{EMG}}$; Fig. 4A; Table 1) as is widely known in the literature (Conway et al., 1995; Salenius et al., 1996, 1997) but also with the fluctuations of the contraction force (Coh$_{\text{force}}$; Fig. 4B; Table 1). Coh$_{\text{EMG}}$ and Coh$_{\text{force}}$ were statistically significant in all subjects ($p$ values $<0.05$; surrogate-data-based statistics), and their maximum values were highly correlated ($r = 0.90, p < 0.0001$; Spearman correlation). The peak frequencies and maximum values of Coh$_{\text{EMG}}$ and Coh$_{\text{force}}$ did not differ statistically significantly from each other ($p = 0.11$ and $p = 0.068$, respectively; Wilcoxon test). These results are in line with a previous finding that, during static extension of the wrist, SM1 activity is coherent at $20\,\text{Hz}$ with finger vibrations recorded with an accelerometer (Airaksinen et al., 2015).
The existence of ~20 Hz Cohforce implies that steady contractions inherently involve ~20 Hz force fluctuations that are coherent with ~20 Hz SM1 oscillations. A close inspection of the force amplitude spectra (Fig. 4B) reveal clear peaks at ~10 Hz, the typical frequency of physiological tremor (McAuley et al., 1997; Gilbertson et al., 2005), but not at ~20 Hz (for amplitude values, see Table 2). The existence of genuine ~20 Hz tremor has nevertheless been demonstrated previously by showing its synchronization with ~20 Hz MEGSM1 signals (Airaksinen et al., 2015), which was evident also in the current study, and by relating finger ~20 Hz tremor to impairment in motor performance (McAuley et al., 1997; Gilbertson et al., 2005).

A natural question to ask is whether the sensorimotor system is sensitive enough to detect reafferent signals associated with the nonsalient ~20 Hz force fluctuations (Fig. 4B, C). Our response is a definitive yes because the Golgi tendon organs can detect stretches as low as 30–90 μN during active muscle contraction (Binder et al., 1977) and thus are sensitive enough to detect the force fluctuations of 2900 ± 1200 μN we observed in the current study (values integrated over a 5 Hz band from 17.5 to 22.5 Hz). Nevertheless, the directional coupling strength was 5.6 times stronger in the efferent than in the afferent direction, as quantified by rPDC (mean across 10–30 Hz; Fig. 4D; p = 0.00035, Wilcoxon test), implying that ~20 Hz Cohforce mainly reflects effects of the efferent cortical drive, similarly to what has been demonstrated for cortex–muscle coherence (Salenius et al., 1997; Gross et al., 2000; Witham et al., 2011; Lim et al., 2014).

**Link between cortex-muscle coherence and burstiness of brain and peripheral signals**

Here we relate the interindividual variability in the burstiness of brain and peripheral signals (quantified with CoV-E) to that in Cohforce and CohEMG to draw conclusions about the causal influence of ~20 Hz brain and peripheral signals on one another. The CoV-E of MEGSM1 (CoV-E_sm1), EMG (CoV-EEMG), and force (CoV-Eforce) peaked for carrier signal at ~20 Hz, as was observed in all subjects for CoV-E_Sm1 (Figs. 5A, 6A) and in the majority of the subjects for CoV-Eforce (Fig. 5B) and CoV-EEMG (Fig. 6B). Indeed, CoV-E_Sm1, CoV-EEMG, and CoV-Eforce peak values in 10–30 Hz exceeded significantly the ~0.51 level expected for white noise for all subjects (p < 0.05), except for 2 subjects for CoV-EEMG. Importantly, the ~20 Hz peak of individual subjects’ CoV-E_Sm1, CoV-Eforce, and CoV-EEMG tightly matched in all subjects, except in these 2 subjects lacking significant CoV-EEMG therefore supporting physiological rather than artifact origins of these envelope modulations.

Moreover, the spatial pattern of the 20 Hz CoV-E_Sm1 agreed with the origin of the signal variations in the rolandic region (Figs. 5A, 7A). Of note, the corresponding topography obtained with task-free MEG data was similar (Fig. 7B), and so were the
As last confirmation of the physiological origin of ~20 Hz envelope fluctuations, the ~20 Hz bursts in MEGSM1 and peripheral signals occur in synchrony. This can be appreciated from the raw envelope traces displayed in Figure 2D, H, L. More quantitatively, MEGSM1–peripheral envelope correlation peaked for carrier frequencies of ~20 Hz in the majority of our subjects (Fig. 8). This was expected since significant envelope correlation was previously reported for ~20 Hz MEGSM1 and EMG signals (Bayraktaroglu et al., 2013).

Across subjects, the maximum CoV-ESM1 across the 10–30 Hz range correlated with the maximum coherence across the same range (Spearman correlation; Coh\textsubscript{force} = 0.73, p = 0.0012, see Fig. 6A; Coh\textsubscript{EMG}, r = 0.73, p = 0.0012, see Fig. 6A), but the association was not strong (Pearson correlation; Coh\textsubscript{force} = 0.75, p = 0.0008, see Fig. 5A; Coh\textsubscript{ESM1}, r = 0.097).

Figure 7. Sensor topography of the CoV of the envelope of ~20 Hz MEG signals (CoV-EMEG) during the isometric contraction task (A) and during the task-free session (B). Subjects are ordered according to their maximum coherence between MEG and force signals (Coh\textsubscript{force}) across 10–30 Hz.

Figure 8. Envelope squared-correlation of MEG\textsubscript{SM1} with peripheral signals (EMG and force; one trace per subject; n = 17). The correlation was computed between the envelopes of the signals filtered through a 5-Hz-wide bands centered on frequencies from 5 to 40 Hz by steps of 1 Hz.
0.66, \( p = 0.004; \) Coh
\textsubscript{EMG} \( r = 0.61, \) \( p = 0.01 \); some subjects with almost identical CoV-
\textsubscript{EMG} had coherence values differing by a factor of \( \sim10. \) The maximum CoV-
\textsubscript{Eforce} and CoV-
\textsubscript{EMG} across the 10–30 Hz range correlated with the corresponding maximum coherences across the same range (Spearman correlation; Coh
\textsubscript{Eforce} \( r = 0.93, \) \( p < 0.0001; \) Coh
\textsubscript{EMG} \( r = 0.81, \) \( p = 0.0001 \), see Fig. 6B), and the association was strong (Pearson correlation: force, \( r = 0.92, \) \( p < 0.0001; \) EMG, \( r = 0.86, \) \( p < 0.0001 \)). Indeed, the association with the coherence magnitude was significantly stronger with CoV-
\textsubscript{Eforce} than with CoV-
\textsubscript{EMG} \( z = 2.47, \) \( p = 0.013; \) Steiger test), and marginally stronger with Coh
\textsubscript{EMG} than with Coh
\textsubscript{Eforce} \( z = 1.87, \) \( p = 0.061) \).

These results confirm that \( \sim20 \) Hz cortex–muscle coherence is tightly linked to the presence of \( \sim20 \) Hz bursts in MEG
\textsubscript{SM1} and peripheral signals; but ultimately, the periphery appears to be the limiting factor. Indeed, some subjects with elevated magnitude of MEG
\textsubscript{SM1} bursts had low coherence, whereas, in contrast, subjects with low (respectively high) CoV-
\textsubscript{Eforce} had systematically low (respectively high) Coh
\textsubscript{Eforce}. This relationship is illustrated in Figure 9 where two subjects with similar \( \sim20 \) Hz CoV-
\textsubscript{ESM1} have strikingly different \( \sim20 \) Hz Coh
\textsubscript{Eforce} while their CoV-
\textsubscript{Eforce} and Coh
\textsubscript{Eforce} vary hand in hand. We take these findings as evidence that, in \( \sim20 \) Hz cortex–muscle coherence, the bursting SM1 activity drives the periphery and the \( \sim20 \) Hz bursts are transmitted with a subject-dependent efficiency.

<3 Hz coupling with force

Although peaking at \( \sim20 \) Hz, Coh
\textsubscript{Eforce} was also salient <3 Hz (Fig. 4C). This frequency range corresponds to the strongest bulk of force fluctuations. Indeed, the power of force fluctuations is \( \sim35 \) times stronger at 1 Hz than at 20 Hz (Fig. 4G, Table 2). A dedicated coherence analysis revealed that both (1) \( \sim20 \) Hz MEG
\textsubscript{SM1} envelope and (2) MEG
\textsubscript{SM1} signals were significantly coherent with both force (Fig. 10A) and its absolute change rate (Fig. 10B) in 13–16 subjects of 17 (\( p < 0.05; \) mean coherence across 0.5–3 Hz; Table 3).

The MEG signals related to these four investigated couplings originated from the SM1 cortex, as was confirmed by individual coherence maps (Figs. 11, 12) and by group-level source-space maps (Fig. 10A,B). At the SM1 sensor with highest mean coherence, the coherence peaked at frequencies between 0.6 and 2.4 Hz (Fig. 10A,B, Table 3).

Directionality analyses revealed a significantly stronger afferent than efferent coupling (quantified with the mean rPDC across 0.5–3 Hz) between (1) force fluctuations and the \( \sim20 \) Hz MEG
\textsubscript{SM1} envelope (\( p = 0.0023; \) Wilcoxon test; Fig. 10C), and (2) the absolute change rate of the force and the MEG
\textsubscript{SM1} signals (\( p = 0.013; \) Wilcoxon test; Fig. 10D). No statistically significant differences in the mean rPDC were found for the other pairs of signals (\( p \) values >0.05).

We further characterized the dynamics of brain signals with respect to changes in force using the TRFs (Fig. 10E,F). Table 4 shows the percentage of variance explained by TRFs. TRFs of the \( \sim20 \) Hz MEG
\textsubscript{SM1} envelope associated with force consistently decreased within 100 ms following the force signal (group-level minimum at 60 ms) and showed weaker, although still prominent, increase peaking \( \sim200 \) ms before the force signal. Individual TRFs of MEG
\textsubscript{SM1} signals associated with force consistently showed their highest values around time 0, but the phase of the oscillations was inconsistent across subjects. Similar observations can be made on TRFs associated with the absolute change rate of the force, but this time, the maximum decrease of \( \sim20 \) Hz MEG
\textsubscript{SM1} envelope occurred later (peak at 220 ms).

Impact of heartbeats on low-frequency coupling

As the coupling we have just outlined occurs at frequencies overlapping with heart rate and its harmonics, we studied in detail the impact of heartbeats on force and brain signals. Figure 13 presents the TRFs of force, MEG
\textsubscript{SM1}, and \( \sim20 \) Hz MEG
\textsubscript{SM1} envelope associated with heartbeats. Time-locked to magnetocardiographic R peak, subjects’ contraction force consistently reached a minimum at \( \sim200 \) ms and was increased by, on average, 17 mN (which corresponds to a movement of \( \sim1 \) μm) at \( \sim400 \) ms. These timings are consistent with known delays between the electric and ballistocardiographic signals (Kim et al., 2016). No consistent trend was visible on subjects’ MEG
\textsubscript{SM1}, and \( \sim20 \) Hz MEG
\textsubscript{SM1} envelope responses to heartbeats. Overall, the TRFs explained only \( \sim4\% \) of the variance of the 0.5–5 Hz content of force, and even less of that of MEG
\textsubscript{SM1} (\( \sim1\% \)) and \( \sim20 \) Hz MEG
\textsubscript{SM1} envelope (\( \sim0.3\% \); Table 4). Importantly, the proportion of force variance explained by heart pulses was not predictive of the level of 0.5–3 Hz coherence between force and \( \sim20 \) Hz MEG
\textsubscript{SM1} envelope (\( r = 0.16, \) \( p = 0.53; \) Spearman correlation) or between force and MEG
\textsubscript{SM1} signals (\( r = 0.076, \) \( p = 0.77; \) Spearman correlation). Finally, the coherence in the sensorimotor sensors was mainly preserved when coherence between force and brain signals was controlled for heartbeats (Fig. 14, in comparison with Fig. 11). Indeed, at the MEG
\textsubscript{SM1} sensor, the ratio of partial coherence controlled for heartbeats to regular coherence was 0.97 ± 0.12 (mean ± SD; range 0.59–1.08) for coherence between force and \( \sim20 \) Hz MEG
\textsubscript{SM1} envelope, and 0.84 ± 0.20 (mean ± SD; range 0.21–1.02) for coherence between force and MEG
\textsubscript{SM1}. Contrastingly, controlling for heartbeats led to strik-
Force \( \frac{dF}{dt} \) directed coherence (\( rPDC \)).

The mean coherence across 0.5–3 Hz and corresponding peak frequencies (mean, SD, and range across subjects)

<table>
<thead>
<tr>
<th>Coherence strength</th>
<th>0.5–3 Hz coherence</th>
<th>Absolute change rate of the force</th>
</tr>
</thead>
<tbody>
<tr>
<td>( 0.5–3 \text{ Hz} ) envelope</td>
<td>0.20 ( \pm ) 0.017</td>
<td>0.055 ( \pm ) 0.037</td>
</tr>
<tr>
<td>Slow ( \text{MEG}_{20} \text{M1} ) activity</td>
<td>0.19 ( \pm ) 0.008</td>
<td>0.009 ( \pm ) 0.032</td>
</tr>
<tr>
<td>Peak frequency [Hz]</td>
<td>20 Hz ( \text{MEG}_{20} \text{M1} ) envelope</td>
<td>1.27 ( \pm ) 0.41</td>
</tr>
<tr>
<td>Slow ( \text{MEG}_{20} \text{M1} ) activity</td>
<td>1.64 ( \pm ) 0.67</td>
<td>0.8 – 2.6</td>
</tr>
</tbody>
</table>

Values reported here were taken from the gradiometer pair of maximum mean Coh_{force} for each subject separately.

For indicative purpose, an orange dotted line indicates the significance level of the mean coherence across 0.5–3 Hz. The plots display individual TRFs (black traces) as well as the mean TRF across subjects (red trace) and \( \pm 2 \text{ SEM} \) (cyan area). Note that we reversed the polarity of TRFs of \( \text{MEG}_{20} \text{M1} \) signals (when the scalar product with the first principal component of sensor orientation was negative) to ensure that all TRFs were estimated in a compatible orientation. This was necessary to warrant the validity of the across-subjects averaging procedure.

Table 3. Mean Coh_{force} across 0.5–3 Hz and corresponding peak frequencies (mean, SD, and range across subjects)

Figure 10. Coupling of \( \text{MEG}_{20} \text{M1} \) with < 3 Hz force fluctuations. A, Coherence between force and \( \text{MEG}_{20} \text{M1} \) signals (left), and between force and \( \sim 20 \text{ Hz MEG}_{20} \text{M1} \) envelope (right). Sub-blocks display the group-averaged coherence source map at 0.5–3 Hz (top) and the individual coherence spectra with the \( \text{MEG}_{20} \text{M1} \) signals. For indicative purpose, an orange dotted line indicates the significance level of the mean coherence across 0.5–3 Hz. B, Same as in A with the force replaced by its absolute change rate \( \frac{dF}{dt} \). C, D, Directional coupling quantified with renormalized partial directed coherence (\( rPDC \)). E, F, TRFs associated to force (E) and \( \frac{dF}{dt} \) (F) of < 5 Hz \( \text{MEG}_{20} \text{M1} \) (left) and \( \sim 20 \text{ Hz MEG}_{20} \text{M1} \) envelope (right). The plots display individual TRFs (black traces) as well as the mean TRF across subjects (red trace) \( \pm 1 \text{ SEM} \) (blue area) and \( \pm 2 \text{ SEM} \) (cyan area). Note that we reversed the polarity of TRFs of \( \text{MEG}_{20} \text{M1} \) signals (when the scalar product with the first principal component of sensor orientation was negative) to ensure that all TRFs were estimated in a compatible orientation. This was necessary to warrant the validity of the across-subjects averaging procedure.
Because heartbeats consistently modulated force signals, we next tackled the question of whether these modulations would play a prominent physiological role in the 0.5–3 Hz coherence between force and SM1 activity. To that aim, we compared the different coherence estimates at $f_{\text{heart-rate}}$ and $f_{\text{off-heart-rate}}$. The mean coherence between force and $\text{MEG}_{\text{SM1}}$ signals at $f_{\text{heart-rate}}$ ($0.021 \pm 0.009$) was $\sim 20\%$ higher than that at $f_{\text{off-heart-rate}}$ ($0.018 \pm 0.007$), but this difference did not reach statistical significance ($p = 0.08; \text{Wilcoxon test}$). This result is in contrast with the finding that coherence between heartbeat and $\text{MEG}_{\text{SM1}}$ signals was $\sim 3$ times higher at $f_{\text{heart-rate}}$ ($0.025 \pm 0.025$) than at $f_{\text{off-heart-rate}}$ ($0.008 \pm 0.010; p = 0.0006$) and that the coherence between heartbeat and force signals was $\sim 4$ times higher at $f_{\text{heart-rate}}$ ($0.049 \pm 0.049$) than at $f_{\text{off-heart-rate}}$ ($0.013 \pm 0.015; p = 0.0003$). Hence, we can conclude that, beyond heartbeats, there exists a prominent and independent coupling between force and SM1 activity at 0.5–3 Hz.

**Discussion**

Our data favor the view that $\sim 20$ Hz cortex–muscle coherence observed during isometric contraction builds on the presence of $\sim 20$ Hz SM1 oscillations and needs not rely on feedback from the periphery. More importantly, our data suggest that effective cortical proprioceptive processing operates at lower, $<3$ Hz frequencies, even when the motor task does not involve voluntary movements. In other words, during steady isometric contraction, small fluctuations in muscle contraction are processed similarly as larger fluctuations occurring during continuous movements (Kelso et al., 1998; O’Suilleabhain et al., 1999; Jerbi et al., 2007; Bourguignon et al., 2011, 2012; Piitulainen et al., 2013a, b) and slow tracking movements (Dipietro et al., 2011; Hall et al., 2014).

**Pervasiveness of $<3$ Hz SM1 fluctuations**

Based on our findings, we suggest the following mechanism for the maintenance of stable muscle contraction: The summed ac-
The activity of proprioceptors sensitive to the force or its change rate (possibly in comparison with expected proprioceptive feedback) (Scott, 2012) modulates population-level SM1 excitability and the 20 Hz SM1 oscillations to compensate their cause. This homeostatic mechanism would directly account for how the brain gets informed about the low-frequency (~3 Hz) content of force fluctuations, which is the strongest and most relevant feedback signal, and needs to be continuously regulated to maintain a stable contraction. This interpretation emphasizes the importance of somatosensory feedback to control motor actions, in line with the influential view that the brain acts as a feedback controller for motor actions (Todorov and Jordan, 2002; Scott, 2004, 2012). Moreover, our finding sheds light on the type of signals the brain relies on to operate the way feedback controllers do.

Our MEG results do not tell whether increases and decreases of force were processed by identical, overlapping, or distinct neu-
SM1 signals are also triggered by the proprioceptive information (Poli et al., 2011), which explains that the SM1 signals change rate of the force rather than by the force itself is in line with generalizes the previous observation that SM1 signals were driven by the absolute activity impacts contraction force during weak isometric finger contractions in the 0–6 Hz range among other frequencies (Sosnoff et al., 2011).

Still, heartbeats seemed to have only a limited impact on the coupling between <3 Hz force fluctuations and MEG signals. Indeed, using partial coherence to control for ECG signals attenuated the coupling with force at sensorimotor sensors by <20% while it effectively removed heartbeat-related artifacts in the lowest sensors of the helmet; these planar sensors are typically the most contaminated by the cardiac artifacts (Jousmäki and Hari, 1996). Moreover, coherence at f_{off-heart-rate} was <20% lower than coherence at f_{heart-rate}. These converging values indicate that heartbeats contributed to <20% of the magnitude of the coupling between <3 Hz force fluctuations and MEG signals. This percentage of coherence relates to both heartbeat artifacts in MEG signals and genuine physiological modulation of the force by the heartbeats.

**Generation mechanism for the ~20 Hz cortex–muscle coherence**

Previous studies have suggested a central role for the ~20 Hz SM1 oscillations in both encoding the motor command and processing the resulting proprioceptive feedback (Riddle and Baker, 2005; Baker, 2007; Witham et al., 2011; Aumann and Prut, 2015). Two action mechanisms have been hypothesized. First, the ~20 Hz cortex–muscle coherence could arise because the SM1 cortex sends pulsed output and monitors the resulting afferent signals to probe the state of the periphery (Mackay, 1997; Baker, 2007; Witham et al., 2011). The second proposed mechanism posits that the ~20 Hz cortex–muscle coherence reflects the integration of afferent information into motor commands to promote a stable motor state (Gilbertson et al., 2005; Androulidakis et al., 2006, 2007; Baker, 2007; Witham et al., 2011). However, our data challenge both these views and suggest that ~20 Hz cortex–muscle coherence builds on the presence of ~20 Hz SM1 oscillations and needs not rely on feedback from the periphery. We therefore propose below a hypothetical generation mechanism of ~20 Hz cortex–muscle coherence.

In monkeys maintaining isometric contraction, the spiking activity of pyramidal-tract neurons is coupled with local field potentials in the SM1 cortex at both ~10 and ~20 Hz (Baker et al., 2003). In other words, the excitability of the pyramidal neurons relates to the phase of the ~10 and ~20 Hz SM1 oscillations; and consequently, the common pyramidal output tends to structure according to these oscillations. However, cortex–muscle coherence seldom peaks at ~10 Hz, which speaks for the existence of a blocking mechanism that prevents the excitability of motoneuron pools from oscillating at ~10 Hz despite the presence of these frequencies in the corticospinal drive (Baker et al., 2003). Such a blocking was argued to be important to prevent excess change. Enhancement of the ~20 Hz SM1 rhythm has been suggested to indicate inhibition or deactivation of the motor cortex (Jasper and Penfield, 1949). The reason why the ~20 Hz SM1 rhythm is modulated with such dynamics can only be speculated. The possible associated effects include transient loss of fine-motor control (e.g., due to attentional reallocation), which generates proprioceptive feedback and/or correction of the ongoing motor plan.

**Impact of heartbeats on force and brain signals**

Heartbeats were associated with force changes of ~17 mN, which explained ~4% of the variance of slow force fluctuations (0.5–5 Hz). This finding is in line with a previous report that cardiovascular activity impacts contraction force during weak isometric finger contractions in the 0–6 Hz range among other frequencies (Sosnoff et al., 2011).

The possible associated effects include transient loss of fine-motor control (e.g., due to attentional reallocation), which generates proprioceptive feedback and/or correction of the ongoing motor plan.
~10 Hz physiological tremor (Baker et al., 2003). Given the close-to-harmonic relationship between ~10 and ~20 Hz SM1 oscillations, we simply suggest that the same mechanism tends to block ~20 Hz oscillatory motor output as well. The ~20 Hz cortex–muscle coherence would then emerge due to nonperfect blocking of the ~20 Hz oscillatory input to the spinal α motoneurons.

**Perspectives for the ~20 Hz cortex–muscle coherence**

Abnormally high ~20 Hz cortex–muscle coherence has been observed in patients suffering from dysfunction of the sensorimotor system, especially of myoclonic disorders (Timmermann et al., 2001; Raethjen et al., 2002; Sileñ et al., 2002; Caviness et al., 2003; Grosse et al., 2003; Kristeva et al., 2004). Given the very strong association between the magnitude of cortex–muscle coherence and the burstiness of ~20 Hz force or EMG signals, we suggest that the observed abnormally high cortex–muscle coherence primarily reflects altered functioning at the level of the periphery, which would be seen (but has not yet been addressed) as high burstiness of both force and EMG due to the myoclonic disorder. Thus, a future alternative means to estimate the level of cortex–muscle coupling would be to quantify the burstiness (with the CoV-E) of both force and EMG; these signals are indeed easily accessible, as they only require measurement of contraction force or muscle activity, with no need to record brain activity.

Finally, our findings provide a novel insight into a so far unresolved puzzle about the cortex–muscle coherence, that is, its great interindividual variability, even in healthy subjects. It was reported that the maximum level of cortex–muscle coherence correlates positively with force CoV and force power in the α and β bands, but the Pearson correlation coefficients were <0.65 (Ushiyama et al., 2017). Here, we demonstrate a positive corre-
luation between maximal cortex–muscle coherence and CoV-E of ~20 Hz force and EMG fluctuations characterized by Pearson correlation coefficients of ~0.9. Accordingly, the present study considerably clarifies the situation by revealing that individual levels of ~20 Hz cortex–muscle coherence are strongly related to the “burstiness” of EMG and force signals at ~20 Hz. In other words, the questions “why is the ~20 Hz cortex-muscle coherence fraught with so high interindividual variability” and “why does the sensorimotor system of different individuals suppress differently the ~20 Hz content of the motor command” are equivalent.

References
Kristeva R, Patino L, Omlor W (2007) Beta-range cortical motor spectral power and corticomuscular coherence as a mechanism for effective cor-