Bodily rhythms appear as novel scaffolding mechanisms orchestrating the spatiotemporal organization of spontaneous brain activity. Here, we follow-up on the discovery of the gastric resting-state network (Rebollo et al., 2018), composed of brain regions in which the fMRI signal is phase-synchronized to the slow (0.05 Hz) electrical rhythm of the stomach. Using a larger sample size ($n = 63$ human participants, both genders), we further characterize the anatomy and effect sizes of gastric-brain coupling across resting-state networks, a fine grained cortical parcellation, as well as along the main gradients of cortical organization. Most (67%) of the gastric network is included in the somato-motor-auditory (38%) and visual (29%) resting state networks (RSNs). Gastric brain coupling also occurs in the granular insula and, to a lesser extent, in the piriform cortex. Thus, all sensory and motor cortices corresponding to both exteroceptive and interoceptive modalities are coupled to the gastric rhythm during rest. Conversely, little gastric-brain coupling occurs in cognitive networks and transmodal regions. These results suggest not only that gastric rhythm and sensory-motor processes are likely to interact, but also that gastric-brain coupling might be a mechanism of sensory and motor integration that mostly bypasses cognition, complementing the classical hierarchical organization of the human brain.

Key words: autonomic; cortical gradients; electrogastrogram; gastric; phase synchronization; resting state networks

**Significance Statement**

While there is growing interest for brain-body communication in general and brain-viscera communication in particular, little is known about how the brain interacts with the gastric rhythm, the slow electrical rhythm continuously produced in the stomach. Here, we show in human participants at rest that the gastric network, composed of brain regions synchronized with delays to the gastric rhythm, includes all motor and sensory (vision, audition, touch and interoception, olfaction) regions, but only few of the transmodal regions associated with higher-level cognition. Such results prompt for a reconsideration of the classical view of cortical organization, where the different sensory modalities are considered as relatively independent modules.
The division of cortical regions into sensory-motor regions, required for the interaction with the external environment, and transmodal regions, devoted to higher-level cognition, is a recurrent feature not only in the resting-state network literature but also more generally in influential proposals of brain hierarchical organization (Fellows and Van Essen, 1991; Mesulam, 1998; Markov et al., 2013). The organization of the brain in relatively independent sensory and motor modules coordinated by higher-order, transmodal regions leaves little space for the processing of internal bodily information. Despite mounting evidence that bodily rhythms and brain activity at rest are tightly coupled (Tort et al., 2018; Azzalin et al., 2019), RSNs are rarely related to internal bodily information (for notable exceptions, see Chang et al., 2013; Shokri-Kojori et al., 2018). Recently, we reported the existence of coupling between brain activity at rest and the slow rhythm generated in the stomach (Richter et al., 2017; Rebollo et al., 2018), a finding which was replicated by an independent research group in a single participant scanned multiple times (Choe et al., 2020). The gastric rhythm is a slow (0.05 Hz) electrical oscillation that is intrinsically generated in the stomach wall by a specialized cell type, known as interstitial cells of Cajal (Sanders et al., 2014), and that can be measured noninvasively with cutaneous abdominal electrodes (Koch and Stern, 2004; Wolpert et al., 2020). The gastric rhythm is produced at all times, but during digestion, the amplitude of the rhythm increases through modulatory influences of the autonomous nervous system (Travaglì et al., 2006; Cruz et al., 2007; Sveshnikov et al., 2012), setting the pace for the contraction of smooth muscles necessary to grind and mix ingested material and eject it into the small intestine.

In the present study, we present a follow-up on the gastric network, a novel RSN composed of brain regions whose fMRI signals are phase synchronized to the gastric rhythm during resting state fixation (Rebollo et al., 2018), in participants who were not in the active phase of digestion. We used a larger sample size ($n = 63$) to provide a detailed characterization of the anatomic extent of the human gastric network across the canonical seven RSNs (Yeo et al., 2011) and across regions of a recent fine-grained multimodal parcellation of the cerebral cortex (Glasser et al., 2016). We additionally computed effect sizes and compared them across regions, which could in turn guide sample size estimation for future studies.

We also quantified how the gastric network was positioned along the main two gradients of cortical organization (Margulies et al., 2016). Finally, we explored how different personal, physiological and experimental variables co-varied with the strength of gastric coupling across participants.

Materials and Methods

Participants

Seventy-two right-handed human participants took part in this study. Thirty-four participants took part in our first study (Rebollo et al., 2018, sample one) and an additional 38 participants (sample two) were recruited for the present study. All volunteers were interviewed by a physician to ensure the following inclusion criteria: the absence of digestive, psychiatric or neurologic disorders; body mass index (BMI) between 18 and 25, and compatibility with MRI recordings. Participants received a monetary reward and provided written informed consent for participation in the experiment and publication of group data. The study was approved by the ethics committee Comité de Protection des Personnes–Ile de France III (approval identifier: 2007-A01125-48). All participants were instructed to fast for at least 90 min before the recordings. Data from nine participants were excluded. Three were excluded because excessive head movement during acquisition (translations larger than 3 mm or rotations larger than 3°), three were excluded because their electrocardiogram (ECG) spectrum did not show a clear peak that could allow us to identify the frequency of their gastric rhythm, and three more were excluded because $<70\%$ of their ECG cycles was within the normogastric range (15–30 s per cycle; Wolpert et al., 2020). A total of 63 participants (mean age 23.95 ± SD 2.76, 31 females and 32 males, and mean BMI 21 ± SD 1.8) were included in the analysis described below.

MRI data acquisition

MRI was performed at 3 Tesla using a Siemens MAGNETOM Verio scanner (Siemens) with a 32-channel phased-array head coil. The resting-state scan lasted 900 s during which participants were instructed to lay still and fixate on a bull’s eye on a gray background. Functional MRI time series of 450 volumes were acquired with an echoplanar imaging (EPI) sequence and the following acquisition parameters: TR = 2000 ms, TE = 31 ms, isotropic voxel size = 3 × 3 × 3 mm$^3$. Each volume comprised 40 contiguous axial slices covering the entire brain. High-resolution T1-weighted structural MRI scans of the brain were acquired for anatomic reference after the functional sequence using a 3D gradient-echo sequence. The two samples had different anatomic sequences. The acquisition parameters for the anatomic scan of sample one were the following: TE = 1.99 ms, TR = 5000 ms, TI-1 = 700 ms/TI-2 = 2500 ms, flip angle-1 = 4°/flip angle-2 = 5°, bandwidth = 240 Hz/pixel, acquisition matrix = 240 × 256 × 224, and isometric voxel size = 1.0 mm$^3$, total anatomic sequence duration = 11 min and 17 s. The acquisition parameters of sample two were the following: TE = 3.24 ms, TR = 2300 ms, TI-1 = 900 ms/flip angle-1 = 9°, bandwidth = 210 Hz/pixel, acquisition matrix = 224 × 256 × 256, and isometric voxel size = 1.0 mm$^3$, with a total anatomic sequence duration of 5 min and 21 s.

Physiologic signal acquisition

Physiologic signals were simultaneously recorded during functional MRI acquisition using MRI compatible equipment and the same montage as in Rebollo et al. (2018). Briefly, physiological signals were recorded 30 s before and after the resting state acquisition to avoid the spreading of the ringing artifact caused by the start of the MRI acquisition. The EGG and electrocardiogram (ECG) were acquired using bipolar EMG electrodes (20 kΩ, 120 cm long) connected to a BrainAmp amplifier (Brain Products) placed between the legs of participants; the amplifier received a trigger signaling the beginning of each MRI volume. EGG was acquired at a sampling rate of 5000 Hz and a resolution of 0.5 μV/bit with a low-pass filter of 1000 Hz and no high-pass filter (DC recordings). ECG was acquired at a sampling rate of 5000 Hz and a resolution of 10 μV/bit with a low-pass filter of 1000 Hz and a high-pass filter of 0.016 Hz. Before the recordings, the skin of participants was rubbed and cleaned with alcohol to remove dead skin, and electrolyte gel was applied to improve the signal-to-noise ratio. The EGG was recorded via four bipolar electrodes placed in three rows over the abdomen, with the negative derivation placed 4 cm to the left of the positive one. The midpoint between the xiphoid process and umbilicus was identified, and the first electrode pair was set 2 cm below this area, with the negative derivation set at the point below the rib cage closest to the left mid-clavicular line. The second electrode pair was set 2 cm above the umbilicus and aligned with the first electrode pair. The third pair was set in the center of the square formed by electrode pairs one and two. The positive derivation of the fourth electrode pair was centered on the
line traversing the xiphoi process and umbilicus at the same level as the third electrode. The ground electrode was placed on the left iliac crest. The ECG was acquired using three bipolar electrodes sharing the same negative derivation, set at the third intercostal space. The positive derivations were set at the fifth intercostal space and separated by 4 cm.

MRI preprocessing
We used the same MRI preprocessing pipeline as described in Rebollo et al. (2018). Brain imaging data were preprocessed using MATLAB (MATLAB 2017, MathWorks) and the Statistical Parametric Mapping toolbox (SPM 8, Welcom Department of Imaging Neuroscience, University College London, United Kingdom). Images of each participant were corrected for slice timing and motion with six movement parameters (three rotations and three translations). Each participant’s structural image was normalized to Montreal Neurologic Institute (MNI) template provided by SPM with affine registration followed by nonlinear transformation (Friston et al., 1995; Ashburner and Friston, 1999). The normalization parameters determined for the structural volume were then applied to the corresponding functional images. The functional volumes were spatially smoothed with a 3 mm3 full-width half-maximum (FWHM) Gaussian kernel.

The time series of voxels inside the brain, as determined using a SPM a priori mask, were subjected to the following preprocessing steps using the FieldTrip toolbox (Oostenveld et al., 2011; Donders Institute for Brain, Cognition and Behavior, Radboud University Nijmegen, the Netherlands; see http://www.ru.nl/brainimaging/fieldtrip, release June 11, 2017). Linear and quadratic trends from each voxel time series were removed by fitting and regressing basis functions, and the blood oxygenation level-dependent (BOLD) time series were then bandpass filtered between 0.01 and 0.1 Hz using a fourth-order Butterworth infinite impulse response filter. A correction for cerebrospinal fluid motion was obtained by regressing out the time series of a 9-mm diameter sphere located in the fourth ventricle (MNI coordinates of the center of the sphere [0–46 –32]).

EGG preprocessing
Data analysis was performed using the FieldTrip toolbox. Data were low-pass filtered below 5 Hz to avoid aliasing and down-sampled from 5000 to 10 Hz. In order to identify the EGG peak frequency (0.033–0.066 Hz) of each participant, we first computed the spectral density estimate at each EGG channel over the 900 s of the EGG signal using Welch’s method on 200-s time windows with 150-s overlap. Spectral peak identification was based on the following criteria: peak power larger than 15 μV2 and sharpness of the peak. Data from the selected EGG channel were then bandpass filtered to isolate the signal related to gastric basal rhythm (linear phase finite impulse response filter, FIR, designed with MATLAB function FIR2, centered at EGG peak frequency, filter width ±0.015 Hz, filter order of 5). Data were filtered in the forward and backward directions to avoid phase distortions and down-sampled to the sampling rate of the BOLD acquisition (0.5 Hz). Filtered data included 30 s before and after the beginning and end of MRI data acquisition to minimize ringing effects.

Heart-rate variability (HRV) preprocessing
In order to computer the power and ratio of HRV, we first removed the MRI gradient artifact from the ECG data using the FMRIB plug-in (Iannetti et al., 2005; Niazy et al., 2005, version 1.21) for EEGLAB (Delorme and Makeig, 2004, version 14.1.1), provided by the University of Oxford Center for Functional MRI of the Brain (FMRIB). Data from the ECG channels were then bandpass filtered (1–100 Hz) using a FIR filter, designed with MATLAB function firw. We then retrieved the interbeat interval (IBI) time series by identifying R peaks using a custom semi-automatic algorithm, which combined automatic template matching with manual selection of R peaks for extreme IBIs. Data from eleven participants was discarded at this stage because of noisy ECG recordings. The resulting IBI time series from the remaining 52 participants were then interpolated at 1 Hz using a spline function (order 3), and the average power in the low (0.06–0.15), and high (0.16–0.4) frequency bands was obtained by means of the Fourier transform using Welch’s method on 120-s time windows with 100-s overlap.

Experimental design and statistical analysis
Quantification of gastric-BOLD phase synchrony
We used the procedure described in Rebollo et al. (2018) to quantify Gastric-BOLD coupling. Briefly, the BOLD signals of all brain voxels were bandpass filtered with the same filter parameters as the ones used for the EGG preprocessing of each participant. The first and last 15 volumes (30 s) were discarded from both the BOLD and EGG time series. The updated duration of the fMRI and EGG signals in which the rest of the analysis was performed was 840 s. The Hilbert transform was applied to the BOLD and EGG time series to derive the instantaneous phases of the signals. The phase-locking value (PLV) (Lachaux et al., 1999) was computed as the absolute value of the time average difference in the angle between the phases of the EGG and each voxel across time (Eq. 1):

$$PLV_{xy} = \left| \frac{1}{T} \sum_{t=1}^{T} e^{i(\phi(t) - \phi(t)))} \right|$$

where T is the number of time samples, and x and y are brain and gastric time series.

Statistical procedure for determining regions showing significant gastric-BOLD coupling at the group level
We employed a two-step statistical procedure adapted from previous work (Richter et al., 2017; Rebollo et al., 2018) to determine which voxels are significantly coupled to the stomach at the group level. We first estimated chance-level gastric-BOLD coupling at each voxel and in each participant. To estimate chance-level gastric-BOLD coupling, we computed gastric-BOLD coupling at each voxel between the BOLD data of the participant and the EGG data of the other 62 participants. For each participant and voxel, chance-level PLV was defined as the median of the 62 surrogate PLVs (Rebollo et al., 2018). Since only the empirical PLVs are specific to the frequency and phase of each participant gastric rhythm, our estimate of chance control for biases in PLV that could be expected using physiological signals with the same length, sampling rate, and frequency range as the empirical EGG, but which are not specific to the exact gastric frequency and phase of that participant. In the second step, we used group-level statistics to determine regions in which gastric-BOLD coupling was greater than chance. To compute group-level statistics, empirical and chance-level PLVs were then compared using a cluster-based statistical procedure that intrinsically corrects for multiple comparisons (Maris and Oostenveld, 2007) implemented in FieldTrip. The procedure consists of applying t tests between empirical PLV and chance-level PLV across participants at each voxel. Candidate clusters are formed by neighboring voxels exceeding a first-level t threshold of p < 0.005 (one-sided). Each candidate cluster is characterized by the sum of the t values in the voxels defining the cluster. To determine the sum of t values that could be obtained by chance, we computed a cluster statistics distribution under the null hypothesis by randomly shuffling the labels “empirical” and “chance level” 1000 times and applying the clustering procedure. At each permutation, we retained the largest positive and smallest negative summary statistics obtained by chance across all voxels and thus built a distribution of cluster statistics under the null hypothesis, and then assessed the empirical clusters for significance. Because the maximal values across the whole brain are retained to build the distribution under the null hypothesis, this method intrinsically corrects for multiple comparisons. Clusters with a Monte Carlo p < 0.025 (one-sided, corrected for multiple comparisons) were considered significant and are reported in Results.

Quantification of gastric-bold shared variance
To estimate the amount of variance in the BOLD signal that could be accounted for by gastric coupling, we computed, for each resting-state network, the squared coherence coefficient between the EGG and average BOLD time course across all voxels significantly coupled to the gastric rhythm using FieldTrip software. The coherence coefficient measures phase and amplitude consistency across time and is a
frequency domain analog of the cross-correlation coefficient in the temporal domain. Therefore, its squared value can be interpreted as the amount of shared variance between two signals at a certain frequency (Bastos and Schoffelen, 2016).

Anatomical characterization of the gastric network across RSNs, cortical gradients, and individual regions

First, we calculated the effect sizes across significant gastric network voxels by computing Cohen’s d (Eq. 2) on difference between empirical and chance PLV, and then projected the resulting volume in MNI space to FreeSurfer’s average template (Desikan et al., 2006) using registration fusion (Wu et al., 2018):

\[ \text{Cohen's } d = \frac{t_{\text{empirical vs chance}}}{\sqrt{n}} \]  

where \( t_{\text{empirical vs chance}} \) is the t statistic of the paired t test between empirical and chance level PLVs, and \( n \) the number of participants.

We then used the seven RSN cortical parcellation available in FreeSurfer native space (Yeo et al., 2011) to identify the networks with more gastric network vertices and with the largest effect sizes. Variability in effect sizes was obtained via bootstrapping. We randomly picked 63 participants with replacement from our original sample, computed Cohen's d's and obtained the standard deviation of this metric across 1000 permutations.

In order to identify the individual regions comprising the gastric network, we used a recent multimodal parcellation of the cerebral cortex (Glasser et al., 2016). This parcellation consists of 180 areas per hemisphere and was obtained using a semi-automated approach in multimodal MRI data that detects sharp changes in cortical thickness, myelin, connectivity, and function. The parcellation was imported to FreeSurfer using the procedure and data available at https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/HCP-MMP1_0_projected_on_Fsaverage/3498446/2. For each region of the parcellation, we determined the overlap with the gastric network (percentage of the area in the gastric network), as well as the average effect sizes across significant gastric network vertices.

We then compared the spatial layout of the gastric network with the first two gradients of functional connectivity described in (Margulies et al., 2016), which we retrieved from neurovault (https://identifiers.org/neurovault.collection:1598). We first resampled the volume containing gastric network significant voxels to match the voxel size of the gradients downloaded from Neurovault (3–2 mm³), using the tool imcalc from SPM. We then divided each gradient into one hundred equidistant bins and quantified for each bin the percentage of cortical voxels belonging to the gastric network. In order to test whether the gastric network is over-represented in particular portions of each gradient, we created a distribution (\( n=1000 \)) of surrogate gastric networks located randomly across the cortex, and recomputed the overlap with each bin and gradient, obtaining a distribution the gastric network across the two gradients under the null hypothesis.

Variability in gastric coupling across participants and regions

We performed a series of exploratory analyses to test for associations between coupling strength, defined as the difference between empirical and chance level PLVs, and personal (gender, BMI, state anxiety score), physiological (EEG and HRV characteristics), or experimental variables (time of day, elapsed time since last meal, head movement, also known as frame-wise displacement). Frame wise displacement was estimated as previously described (Power et al., 2012). State anxiety was measured using the STAI-B questionnaire (Spielberger et al., 1983). Gender and sample effects were assessed using unpaired samples t test. The influence of the rest of the variables was assessed by separate robust linear regressions using R MASS package (Ripley et al., 2013), which included the covariate and the intercept term. Bayes factor was used to quantify the evidence for H0 relative to H1, with a value <0.33 indicating substantial evidence for a null effect (Kass and Raftery, 1995; Wetzels and Wagenmakers, 2012). For the unpaired samples t test, we used the methods described previously (Rouder and Morey, 2011) to determine the Bayes factor. For the regression analysis, an approximation of the Bayes factor was computed (Wagenmakers, 2007) by comparing the Bayesian Information Criterion of the regression model including the covariate and the intercept to a regression model with only the intercept. Group level GLMs on coupling strength across voxels were performed using SPM 12 (Wellcome Department of Imaging Neuroscience, University College London). All variables were mean centered, and the intercept term was included.

Correction of MRI images for cardiac timing

As a control analysis, we computed the spatial extent of the gastric network and gastric coupling strength, with and without adjusting fMRI images for the cardiac cycle using retrospective image correction RETROICOR (Glover et al., 2000). We used the PhysIO toolbox implementation (Kasper et al., 2017) of the RETROICOR algorithm to generate six cardiac phase regressors from the cardiac timings in the 52 participants for whom we had cardiac peak timings. We then regressed out the cardiac regressors together with CSF timeseries, and obtained a new set of empirical and surrogate PLVs as well as a new significant group level network (cluster forming threshold \( p = 0.005 \) one-sided, cluster significance threshold \( p = 0.025 \), one-sided). We then compared the resulting gastric network with that of the same subsample but without RETROICOR correction, and calculated the distribution of the gastric network across Yeo seven networks (Yeo et al., 2011). Finally, we compared how the RETROICOR pipeline affected coupling strength (empirical = surrogate PLV) in the original gastric network (\( n=63 \)), for the 52 participants with and without RETROICOR.

Code and overlays availability

The code necessary to reproduce the results is available at https://github.com/rebollo/StomachBrain_2021. Unthresholded t-maps, effect sizes and significant voxels can be found at https://identifiers.org/neurovault.collection:9985.

Visualization tools

The distribution of coupling strength was plotted using raincloud plots (Allen et al., 2019). Visualization of brain data on the cortical surface was done using PySurfer (https://pysurfer.github.io/).

Results

The stomach is synchronized with somatosensory, motor, visual, and auditory regions

We first determined the frequency of each participant gastric rhythm, by identifying the EGG channel displaying the largest peak within normogastric range (0.033–0.066 Hz). The mean EGG peak frequency across the 63 participants was 0.049 Hz (±SD 0.004, range 0.041–0.057). We did not find statistically significant differences between the two samples in EGG frequency [unpaired samples t test, mean sample one = 0.047 ± 0.003 Hz (±SD), mean sample two = 0.048 ± 0.0036 Hz, \( t(61) = 1.41, p = 0.161 \) Bayes factor = 0.6, indicating not enough evidence to disentangle between H1 and H0] nor in EGG power (unpaired samples t test, mean sample one = 220 ± 410 μV², mean sample two = 333 ± 530 μV², \( t(61) = 1.41, p = 0.356 \), Bayes factor = 0.37, indicating not enough evidence to disentangle between H1 and H0). We did find differences in the standard deviation of EGG cycle length, which was significantly smaller in sample two than sample one (unpaired samples t test, mean sample one = 3.38 ± 1.25 s, mean sample two = 2.66 ± 1.48 s, \( t(61) = -2.08, p = 0.041 \)), indicating a more stable gastric rhythm in sample two.

We then quantified the degree of phase synchrony between the EGG signal and BOLD time series filtered around gastric frequency using empirical and chance-level estimates of PLV. PLV (Lachaux et al., 1999) measures the level of phase synchrony, defined as the stability over time of the time lag between two time series, independently of the amplitude of the two time
series. Importantly, PLV is large as long as the time lag is constant, independently from the value of the time lag, and does not provide information about directionality. In each participant and voxel, we computed the empirical PLV between the narrowband BOLD signals and the gastric rhythm. We also estimated the amount of PLV that could be expected by chance in each voxel using the BOLD signal of one participant and the EGG of the other 62 participants, and taking the median of that surrogate distribution as chance-level PLV. The empirical PLVs were then compared with the chance-level PLVs using a cluster-based statistical procedure that intrinsically corrects for multiple comparisons (Maris and Oostenveld, 2007). Significant phase coupling between the EGG and resting-state BOLD time series occurred in 32 clusters (voxel threshold \( p < 0.005 \), one-sided paired \( t \) test between observed and chance PLV; cluster threshold corrected for multiple comparisons, Monte Carlo \( p < 0.025 \) one-sided). Exact \( p \)-values are reported for each cluster in supporting table Extended Data Figure 1-1.

As observed before (Rebollo et al., 2018), the gastric network (Fig. 1A; Extended Data Fig. 1-1) comprises bilateral regions...
along the central, cingulate, and lateral sulci, as well as occipitoparietal-temporal regions and portions of the left striatum (Fig. 1B), bilateral thalamus (Fig. 1C), and cerebellum (Fig. 1D). The distribution of the average gastric network coupling strength (defined as the difference between empirical and chance PLV) across participants (Fig. 1F) ranged from 0.0026 to 0.2096 (mean = 0.0417, STD = 0.0399, median = 0.0319). We found no difference between the two samples in average coupling strength (unpaired sample t test, mean sample one = 0.043 ± 0.044, mean sample two = 0.039 ± 0.031, t(61) = −0.42, p = 0.670, BF = 0.27, indicating substantial evidence for H0).

We then used a well-known parcellation of the cortical surface into seven RSNs (Yeo et al., 2011) to quantify the overlap of the gastric network with canonical RSNs. This analysis (Fig. 1E) confirmed that most of the gastric network (67%) is included in the somato-motor (38%) network, which also includes the auditory cortices, and in the visual (29%) network. Gastric coupling in the somato-motor-auditory network spans somatosensory, motor and auditory regions surrounding the central, lateral and cingulate sulci. Gastric coupling in the visual network spans striate and extrastriate visual regions, and is particularly pronounced in medial and ventral occipital regions. Effect sizes (Cohen’s d of paired t test of empirical against chance PLV) were similar across visual and somato-motor networks (Fig. 1E, right). Effects sizes in the few coupled regions of the default, saliency, control and attention networks were only slightly smaller than those in sensory and motor cortices (Fig. 1E, right). Similarly, the average shared variance, as estimated from squared coherence, between the EGG and BOLD signal in each RSN, was very similar in all RSNs, ranging from 13.8 ± 2.0% in the limbic system to 15.8 ± 2.2% in the default network. Intersubject variability was high in all RSNs, with shared variance in individual participants ranging from below 1% up to above 40% in all RSNs.

The gastric network showed a very limited overlap with non-sensory networks (Fig. 1E, left). The overlap with the default network (9.5%) occurs mostly in one medial node of the default network, the retrosplenial cortex as well as in the lateral node in the superior temporal sulcus, and a small rostral prefrontal region. Only 8.1% of the gastric network is found in the saliency network, sparing core regions of the saliency network such as the anterior insula and the fundus of the dorsal anterior cingulate sulcus. Only very few regions of the gastric network belonged to the control network (4.6%), dorsal attention network (3.6%), or limbic network (0.4%). Subcortical regions (which in this parcellation include thalamus and striatum but not cerebellum), represented 6.8% of the gastric network. Even if the percentage of the gastric network in subcortical regions is small, effect sizes were similar as those in the cortical regions of the gastric network (Fig. 1E). We do not examine thalamus and striatum in more details because the voxel size we used is not well suited for those structures, and the projection of voxels into surfaces is optimized for cortical regions and hence induce deformation on subcortical regions. Finally, 11.9% of the gastric network is found in the cerebellum. Note however that the most ventral part of the cerebellum was not imaged in this study, so this percentage is only an indication of the share of the cerebellum in the gastric network. Within the cerebellum, the gastric network was mostly found in Left crus I (30.1%), Right lobule VI (16.2%), Left crus II (11.9%), Left lobule VI (11.57%), Vernus crus II (10.4%), according to the parcellation from Diedrichsen et al. (2009).

While the gastric network is spatially extended, effect sizes remain moderate. The average Cohen’s d across visual, somatomotor or default network voxels was close to 0.4, raising the issue of replicability and required sample sizes. For the first-level statistical threshold employed in these analysis (cluster forming threshold = 0.005 one-sided), a sample size of at least 77 participants is required to observe an effect of this magnitude 80% of the time, meaning that it is possible that the current study is slightly underpowered. Still, we wanted to verify whether gastric coupling was more prevalent in sensory-motor cortices across the two subsamples that encompass this study. For this, we recomputed the spatial layout and effect sizes of significant gastric brain coupling in each of the two subsamples (29 and 34 participants, respectively), albeit using a more liberal statistical threshold (cluster forming threshold = 0.05 one-sided), which requires a sample size of 41 participants to detect an effect 80% of the time. The spatial layout of the gastric coupling was indeed similar across the two sub-samples (Fig. 1E), with the largest proportion of the gastric network in somato-motor and then visual networks, and more varying proportion of coupling in transmodal networks across the two subsamples. The relative strength of effect sizes was more variable between the two sub samples, with sample 1 having larger effect sizes in visual than somato-motor networks and an opposite pattern in sample 2.

To verify that the dominance of sensory and motor cortices in the gastric network was not related to the level of details of the parcellation, we used a more recent parcellation of the cerebral cortex (Glasser et al., 2016) consisting of 180 areas per hemisphere. The parcellation was obtained using a semi-automatized approach in multimodal MRI data that detects sharp changes in cortical thickness, myelin, connectivity, and function. For each region of the parcellation overlapping with the gastric network, we computed the percentage of the area in the gastric network and the average effect sizes across significant gastric network vertices (Extended Data Fig. 1-2). The 25 regions with the largest effect sizes across both hemispheres (Fig. 1G), were primary motor and somatosensory cortices (regions 4, 3b, 1, and 3a), premotor [6d, supplementary motor and cingulate eye fields (SCEF), 6mp, frontal eye field (FEF)], cingulate motor (24dd), auditory (STSa, A5, MBelt, primary auditory cortex (A1), PBelt, RI, STGa), insular granular, opercular (area 43) early visual regions (V1, V2, V3 and VVC), area FEF from the saliency/attentional networks, and the retrosplenial complex (RSC), area STSa and area 10d, three regions typically associated with the default network.

Because the whole-brain analysis indicates that all sensory cortices are coupled to the gastric network except the olfactory cortex, we performed a targeted region of interest analysis of the left and right piriform cortex as defined in the Glasser et al. (2016) parcellation. We found that coupling in the olfactory cortex has indeed a medium effect size (Cohen’s d left = 0.56, right = 0.61), comparable to the effect sizes in primary visual (left = 0.57, right = 0.54) or auditory (left = 0.67, right = 0.61) cortices. The absence of olfactory cortices from the whole-brain analysis is thus probably because of the small size of those cortices.

**Gastric network and cortical gradients of functional connectivity**

The gastric network is mostly found in somato-motor-auditory and visual RSNs. To verify that this is not because of the specific resting-state network parcellations we used, or to a somewhat arbitrary division between sensory and transmodal areas, we analyzed how the gastric network is distributed along the first two gradients of functional connectivity described by (Margulies et al., 2016). In this approach, each cortical voxel can be defined by...
its location along two different axes, one that goes from unimodal to transmodal regions, and another one that goes from visual to transmodal to somato-motor-auditory regions. In Figure 2A, we reproduce the findings of Margulies et al. (2016). When considering the whole cortex, the projection of the probability density on the first gradient shows two prominent peaks at the extremities, corresponding to transmodal and unimodal regions (Fig. 2A, red curve). When considering only the gastric network, the probability density shows a markedly different profile, with an increase in the unimodal extreme only (Fig. 2B, red curve). As described by Margulies et al. (2016), the projection of the whole-brain probability density on the second gradient shows a prominent peak in transmodal regions (Fig. 2A, blue curve). However, gastric network voxels are more densely represented in the visual and somato-motor-auditory extremes of the gradient (Fig. 2B, blue curve), indicating that coupling with the gastric rhythm is more likely to be present in unimodal than in transmodal brain regions.

We then tested whether the distribution of gastric network voxels along the first and second gradients could be because of chance. We first computed the percentage of brain voxels that belong to the gastric network along one hundred equidistant bins (Fig. 2C, orange lines). In order to test whether spatial biases of such size could be obtained by chance, the spatial position of the gastric network voxels was randomly permuted across the cortex, while keeping the spatial layout of the gradients intact, and computed the percentage of voxels belonging to the surrogate gastric network thus created. The procedure was repeated one thousand times to estimate the distribution under the null hypothesis (Fig. 2C, black circles). For the first gradient, the percentage of all brain voxels overlapping with the gastric network across bins is systematically larger than chance in the unimodal bins of the gradient, and systematically smaller in the remaining bins (Fig. 2C, left). Similarly, for the second gradient, the overlap with the gastric network across bins is systematically larger than chance in the unimodal bins and systematically smaller in the transmodal bins (Fig. 2C, right). This analysis confirmed that the spatial biases in the unimodal extremes of the gradients could not be obtained by chance.

**Anatomical characterization and effect sizes in the gastric network**

We now examine in more detail the anatomy of the gastric network, starting with regions belonging to the somato-motor network in central and mid-cingulate regions, followed by a characterization of opercular regions in and around the lateral sulci, including auditory cortices and insula, then regions located in the posterior portion of the medial wall and in occipital, posterior cingulate, temporal and parietal cortices, and ending with the few transmodal regions in frontal, prefrontal, and lateral temporal cortices.

---

**Figure 2.** The gastric network and cortical gradients of functional connectivity. A. Density plot depicting the distribution of all cortical voxels along the first two gradients of functional connectivity described previously (Margulies et al., 2016). The first gradient (y-axis) runs from unimodal (negative values, sensory or motor regions) to transmodal regions (positive values). The second gradient (x-axis) runs from visual (positive values) to somato-motor and auditory regions (negative values). The color scale depicts the logarithm of the number of voxels. The projection of the probability density on the first and second gradients are shown in red and blue, respectively. B. Density plot of gastric network voxels on the first two gradients of functional connectivity. Gastric network voxels are located in the unimodal extremes of the two gradients. The projection of the probability density on the first and second gradients are shown in red and blue, respectively. C. Percentage of all brain voxels that belong to the gastric network (orange) for each of the two gradients, computed on 100 equidistant bins. The black circles depict the distribution of chance level overlap obtained by reallocating randomly the spatial position of gastric network voxels in the cortex. Gradients downloaded from https://identifiers.org/neurovault.collection:1598.
**Gastric network surrounding the central and cingulate sulci**

Gastric network in central and mid-cingulate regions in central and mid-cingulate regions (Fig. 3A), the gastric network covers primary somatosensory (green), motor, premotor, and cingulate motor (blue) and nonmotor (violet) cingulate regions, as well as area 5lb (pink). The primary somatosensory cortex proper (areas 3b and 1), is located in the postcentral gyrus and is involved not only in the representation of bodily surface but also of the visceras (Amassian, 1951; Dum et al., 2009; Kern et al., 2013; Azzalini et al., 2019). Accessory somatosensory area 2, in the bank of the postcentral sulcus, shows little coupling with the gastric rhythm. In the anterior direction, area 3a, in the fundus of the central sulcus, is sometimes referred to as part of the somatosensory cortex but can also be seen as a transition zone with primary motor area 4 (Catani, 2017) in the posterior bank of the postcentral gyrus. Besides its role in controlling skeletal muscle, area four has been shown to provide sympathetic input to the stomach (Levinthal and Strick, 2020). Both area 3a and area four display coupling with the gastric rhythm. The gastric network is also found in numerous premotor areas, generally involved in integrating sensory and motor information for the performance of actions: area 6 (6m, 6d, 6v, 6a), the FEF and supplementary motor areas (6mp, 6ma, SCEF). Note that area SCEF extends into the dorsal bank of the anterior cingulate sulcus. Along the medial wall, the gastric network overlaps with left area 5l (cyan), the cingulate motor areas (24dd, 24dv, blue), known to contain several somatotopic maps (Amiez and Petrides, 2014) and to receive spinothalamic input (Dum et al., 2009), the paralimbic cortex (area p32pr), areas p24pr and 23d, located adjacent to the corpus callosum and area 23c, located between the cingulate motor regions and the task-negative precuneus. Finally, the gastric network also includes area 5lb mostly on the right side, a recently discovered region which is left-lateralized for language and right-lateralized for theory of mind (Glasser et al., 2016), and to a lesser extent, the superior frontal language area (SFL), a region that is also left-lateralized for language.

Regions containing a large proportion of voxels belonging to the gastric network (Fig. 3B) include primary somatosensory (3b, 3a, 1), primary motor (4), premotor (SCEF, 6d, FEF), cingulate motor regions (24dd), region 5lb and area 5m. Gastric-brain coupling show a strong right lateralization in regions surrounding the cingulate sulcus (p32pr, p24pr, 23cc 23d, 24dv, 24dd, SCEF), as well as in FEF and area 5lb, while areas 1 and 5m are left-lateralized. Effect sizes (Fig. 3C) were largest in somatosensory (3b, 1, 3a) and primary motor (4) cortices, followed by premotor and cingulate motor regions (6d, SCEF, 24dd, FEF, 6m). Asymmetries in effect sizes showed similarities with asymmetries in overlap, with right lateralization in effect size in cingulate regions (p24pr, 23cc 23d, 24dv, 24dd, SCEF) and region 5lb, and left lateralization in regions 5l, 5m, and 2.

**Figure 3.** Gastric network in central and mid-cingulate regions. A, Effect sizes in the gastric network in central (top) and mid-cingulate regions (bottom) displayed on the left and right inflated surfaces and overlaid with the corresponding regions of Glasser et al. (2016) parcellation. Green, primary somatosensory; blue, motor and premotor regions; pink, area 5lb; violet, nonmotor cingulate regions; cyan, area 5 and its subdivisions. B, Percentage of each region overlapping with the gastric network. C, Effect sizes of gastric-BOLD coupling in voxels belonging to the gastric network split by regions. A, Anterior; I, Inferior; LH, Left hemisphere; P, posterior; RH, Right hemisphere; S, Superior. Area names as in text and Extended Data Figure 1-2.

**Gastric network surrounding the lateral sulcus**

In and around the lateral sulcus (Fig. 4A), the gastric network spans secondary somatosensory (green), frontal operculum (blue), early (dark violet) and association (light violet) auditory regions as well as regions of the insula proper (pink). The secondary somatosensory (SII), known to process inputs from the skin but also from the visceras (Amassian, 1951) is composed of areas OP1, OP2–3, and OP4, and is located anteriorly to area PfcM, in the inferior parietal cortex. SII is separated from the frontal operculum (areas FOP1, FOP2, FOP3, and FOP4) by area 43 (cyan). Early auditory regions, involved in the processing, identification and location of sounds (Jasmin et al., 2019), include the A1 and the surrounding lateral belt (Lbelt), medial belt (MBelt), parabelt (Pbelt), and retro-insular cortex (RI). Auditory association regions extend inferiorly to the superior temporal sulcus and include A4, A5, temporal area 2 (TA2), and auditory default network regions dorsal posterior superior temporal sulcus (STSdp), dorsal anterior superior temporal sulcus (STSda), and ventral anterior superior temporal sulcus (STSva). The insula proper consists of area 52, a transition region between...
auditory cortex and insula, the granular insula (Ig), which contains a complete somatotopic motor map (Glasser et al., 2016) as well as viscerosensitive neurons (Cechetto and Saper, 1987), the middle insular area (MI), and posterior insular area I and II (PoI1, PoI2).

The regions with the largest overlap with the gastric network consist of early auditory regions (Pbelt, A1, MBelt, RI), right insular regions (Ig, 52), left parietal opercular regions (OP4, OP1, 43) and right area FOP1 (Fig. 4B). Effect sizes were largest in early and association auditory regions (A5, A4, MBelt, L Belt, A1, PBelt, STSda, RI, TA2), insular cortex (52, Ig, PoI2), and areas 43 and FOP1 (Fig. 4C). Overall, overlap and effect sizes in SII (OP1, OP2–3) were most prominent in the left hemisphere while overlap and effect sizes in the frontal, insular, MBelt and auditory association regions were strongly right-lateralized.

**Gastric network in posterior regions**

In the posterior part of the brain, the gastric network covers large portions of the occipital visual cortex (Fig. 5A, dark green), occipito-parietal sulcus (blue) and retrosplenial cortex (violet), dorsal precuneus (light green), ventral visual stream (pink), and right temporoparietal-occipital junction (black). The gastric network is found in all early visual regions (V1, V2, V3, and V4), involved in the processing of low-level visual properties such as orientation and color, extending superiority to the dorsal stream (V3A, V3B, V6, V6A, V7), involved in the location of objects in space, up to area 7Am and the precuneus visual area (PCV), a region sensitive to optic flow (Glasser et al., 2016). Medially and anteriorly, the gastric network extends to the parieto-occipital sulcus and recruits portions of the dorsal visual transition area (DVT), prostrate cortex (Pros), and parieto-occipital sulcus regions 1 and 2 (POS1, POS2), regions typically involved in the generation of the electrophysiological α rhythm (Richter et al., 2017) and vigilance fluctuations. Area POS2 stands out as because of its large overlap with the gastric network (90%). Conversely, it is worth noting that the gastric network spares most of the ventral precuneus (yellow; areas 7m, 31pd, 31a, 31pv), which is one of the core nodes of the default network. Ventrally, the gastric network extends to ventromedial visual areas 1, 2, and 3 (VMV1, VMV2, and VMV3), as well as regions of the ventral stream, involved in the processing of objects identity [V8; area lateral occipital 2 (LO2), posterior inferotemporal complex (PIT), and fusiform face complex (FFC)], and the occipital-temporal-parietal junction (Black, MT, Ppg, TOP3, LO3) in the right hemisphere.

A cluster of dorsal right regions (V3A, V6A and V7, POS2, and DVT) stands out, with >80% of those regions in the gastric network (Fig. 5B). Other posterior regions with large overlap were right area VMV3, left V3B, and, to a lesser extent, the RSC, as well as left V1 and ProS. Effect sizes (Fig. 5C) were largest in RSC, right FFC and most occipital visual regions including V1. A rightwards lateralization in effect sizes and overlap was present.
for most regions, with the exception of area V3B, which displayed a leftwards lateralization.

Gastric network in prefrontal and lateral temporal cortex

The gastric network includes a few trans-modal regions located in the middle temporal gyrus (Fig. 6A, pink), inferior frontal cortex (blue), dorsolateral prefrontal cortex (green), and in the orbitofrontal cortex (yellow). In the middle temporal gyrus, the gastric network is found in the middle temporal gyrus (TE1m, TE1p), as well as in ventral regions of the superior temporal sulcus (STSdp, STSvp), belonging to the auditory portion of the default network. The percentage of voxels belonging to those regions remain small (Fig. 6B), with moderate effect sizes (Fig. 6C), and a marked right lateralization.

In right inferior frontal cortex (Fig. 6A, blue), the gastric network forms a cluster that comprises the posterior portion of the inferior frontal sulcus (IFSp), the anterior portion of the inferior frontal junction (IFJa), and area posterior 9–46d (p9–46d) as well as area 44, all regions of the control network, extending to rostral area 6 (6r) of the saliency network. In the left dorsolateral prefrontal cortex (Fig. 6A, green), the gastric network includes multiple regions of the control network, including Superior Transitional Area 6–8 anterior (s6–8a), anterior area 10p (a10p), area 11 lateral (11l), anterior area 9–46 ventral (9–46v), area 9–46 dorsal (9–46d), as well as area 8b lateral from the default network and polar area 10 (10p) from the limbic network. Overlap is particularly pronounced in a10p and a9–46v (Fig. 6B) with an exclusively left lateralization. Finally, in orbitofrontal cortex (Fig. 6A, yellow), the gastric network includes dorsal and rostral portions of area 10 (10d, 10r) from the default.

Variability in gastric coupling across participants and regions

We examined how differences in personal (gender, BMI, state anxiety score), physiological (EEG and HRV characteristics) or experimental variables (time of day, elapsed time since last meal, head movement, also known as frame-wise displacement), were related to variability in coupling strength across participants, in a series of exploratory analyses. Because of the small age range of our sample (18–30), we did not include age as a regressor. We first tested whether any of these variables accounted for a compact measure, the mean coupling strength in the gastric network (Table 1).

We found no differences between genders in the average coupling strength (mean female = 0.039 ± 0.036, mean male = 0.044 ± 0.044, paired t test, t(61) = 0.48, p = 0.634, Bayes factor = 0.279, indicating substantial evidence for the null hypothesis). All other variables were tested with robust linear regressions. With one exception,
none of the variables tested accounted for coupling strength, including state anxiety ($t_{(58)} = 0.47$, $p = 0.634$, Bayes factor = 0.124 indicating substantial evidence for the null hypothesis).

The only variable related to coupling strength was the low-frequency component of HRV (LF-HRV; Fig. 7C), which was associated with a larger brain coupling ($n = 52$, $t_{(50)} = 3.76$, robust regression, $r^2 = 0.22$, uncorrected $p = 0.0004$, Bonferroni corrected $p = 0.0048$). LF-HRV reflects multiple cardiac control mechanisms, including blood pressure and baroreflex activity, and is under the control of both sympathetic and parasympathetic nervous system (Goldstein et al., 2011; Reyes del Paso et al., 2013). Because coupling strength turns out to be related to HRV, we recomputed to the spatial layout of the gastric network with and without correction for cardiac pulsation using RETROICOR (Glover et al., 2000; Kasper et al., 2017), in the 52 participants where heart rate time series were obtained. We observed no overall difference in the distribution of the gastric network across resting networks: gastric-brain coupling was mostly observed in somato-motor-auditory network as well as in the visual network after RETROICOR correction (Fig. 7D). The spatial layout of the gastric network is thus not solely driven by cardiac pulsation effects. We then tested whether correcting for cardiac pulsation effects affected the overall level of coupling strength. We compared coupling strength in the same spatial extent of gastric network (as defined on 63 participants), with and without RETROICOR. Correcting for cardiac pulsation induced a significant reduction in overall coupling strength (Fig. 7E, mean before RETROICOR = $0.043 \pm 0.040$ STD, mean after RETROICOR = $0.026 \pm 0.020$ STD, paired $t$ test $t_{(50)} = 5.42$, $p = 0.000001$). The linear relationship between LF-HRV and coupling strength remained significant after cardiac pulsation correction ($n = 52$, $t_{(50)} = 4.23$, robust regression, $r^2 = 0.26$, $p = 0.00009$).

We reasoned that only some gastric network regions might be modulated by the variables we analyzed. We therefore used two different strategies to look for more subtle effects of interindividual variables on coupling strength, both accounting for multiple comparisons. We first tested for effects of these variables across Glasser et al. (2016) regions of interest (ROIs) overlapping with the gastric network, applying false discovery rate (FDR) correction over the 226 ROIs tested for each variable separately. We only found one significant association, with coupling strength in right area 7Am co-varying with EGG cycle duration variability ($t_{(61)} = 3.936$, FDR $p = 0.048$). We also performed group level general linear models with coupling strength in all brain voxels as the dependent variable. Gender, group, BMI, frame-wise displacement, time of the day, EGG frequency, and EGG power were used in a first general linear model with all participants ($n = 63$). Separate general linear models including these variables and either state anxiety ($n = 61$), elapsed time since last meal ($n = 51$) or HRV measures ($n = 51$) were performed. None of the models survived correction for multiple comparisons (FWE $p < 0.05$).
Discussion

We examined the anatomy of the gastric network, a set of brain regions phase coupled to the rhythm of the stomach during rest, and analyzed the spatial layout of the gastric network at the level of its constituent regions (Glasser et al., 2016), resting-state networks (Yeo et al., 2011), and its extent and position along the first two gradient surfaces of connectome connectivity that underlie the topological structure of the cortex (Margulies et al., 2016). We found that the gastric network is overrepresented in unimodal sensory-motor regions, and underrepresented in transmodal regions. All sensory and motor cortices are coupled to the gastric rhythm, including not only areas responding to touch, vision and audition but also the interoceptive insula and, to a lesser extent, the olfactory piriform cortex. Only few transmodal regions were coupled to the gastric rhythm, mostly in the default network (retrosplenial cortex, right area STSda). We also observed significant coupling in the cerebellum. None of the personal, physiological and experimental variables tested co-varied with the overall strength of gastric coupling across participants, with the exception of an index of cardiac autonomic activity. Notably, we found substantial evidence for an absence of association between state anxiety and gastric-brain coupling during rest.

Anatomical pathways for gastric brain coupling

Some of the gastric network regions we observe are known to be involved in interoception and autonomic functions. Such is the case of the granular insula, the somatosensory cortices, and the cingulate motor regions (Aramassian, 1951; Cecchetto and Saper, 1987; Dunn et al., 2009), which receive visceral inputs, and motor regions, which provide sympathetic input to the stomach (Levinthal and Strick, 2020). However, we find that gastric-brain coupling extends well beyond expected visceral processing regions, notably early and association visual and auditory cortices. Such results are in line with findings in rats, where the electrical stimulation of the stomach elicits BOLD responses not only in somatosensory, insula, cingulate cortices, but also in motor, auditory and visual cortices (Cao et al., 2019). Changes in visual cortices activity has also been observed in humans in response to painful gastric distension (Van Oudenhove et al., 2009), subminal rectal distension (Kern and Shaker, 2002), as well in dogs after gastric electrical stimulation (Yu et al., 2014). Similarly, colonic pain induces a response in the rat auditory cortex (Wang et al., 2008). The parabrachial nuclei, the major relay of both spinal and vagal visceral afferents, projects not only to thalamic relay nuclei involved in interoception and touch (Saper and Loewy, 1980; Coen et al., 2012) but also to the visual lateral geniculate nucleus (Erwig et al., 1997), and potentially to the auditory medial geniculate nuclei (Uhlrich et al., 1988). Projections of the parabrachial nucleus might thus at least partially mediate gastric-BOLD coupling in sensory cortices. It is also worth underlining that we find no gastric-BOLD coupling in anterior insula and medial prefrontal regions, two regions involved in parasympathetic control of the stomach in rats (Levinthal and Strick, 2020).

Beyond direct ascending or descending communication with the stomach, gastric-coupling could also stem from intracortical connectivity. For instance, direct connections have been reported between secondary somatosensory cortex to auditory cortices, between visual area MT and primary somatosensory (Cappe and Barone, 2005). Whether and how our findings pertain to interoceptive prediction models (Barrett and Simmons, 2015; Petzschner et al., 2021) remains an open question, since neither the spatial resolution nor the gradients (Margulies et al., 2016; Huntenburg et al., 2018) we used provide information on the laminar origin of gastric-BOLD coupling. Finally, neuromodulation might also be involved (Rinaman and Schwartz, 2004; Rebollo et al., 2021). In particular, activity in the locus coeruleus, the main source of norepinephrine to the forebrain, is modulated by gastric, colonic and bladder distension (Elam et al., 1986; Saito et al., 2002), inducing fluctuations in arousal that can also be obtained with distension of the small intestine (Kukorelli and Juhász, 1977).

Intersubject variability and cardiac activity

When examining effect sizes and the required sample sizes needed to observe our effect, it appears that the current study might still be slightly underpowered. However, note that we could only estimate the sample size for the first step of our statistical procedure (voxel-level statistics), and not to the second step (cluster-level statistics). A more precise estimation of the required number of participants for future studies remains to be established, considering the intersubject variability observed in gastric-BOLD coupling that was not readily explained by the individual variables we explored.

Indeed, we examined a number of individual variables which could covary with gastric-brain coupling, including state anxiety, which has often been associated with (mostly cardiac)

Table 1. Variables tested to account for average coupling strength in the gastric network

| Variable type | Variable name                           | n   | Min  | Max  | Mean | Median | Std   | t     | p      | BF    |
|---------------|----------------------------------------|-----|------|------|------|--------|-------|-------|--------|-------|
| Personal      | Gender                                 | 63  | 17.5 | 24.5 | 21.0 | 21.0   | 1.8   | 0.52  | 0.604  | 0.0000|
|               | BMI                                    | 63  | 20   | 53   | 32.9 | 31.5   | 8.7   | 0.47  | 0.643  | 0.1240|
|               | State anxiety (STAI B)                 | 63  | 0.061| 0.323| 0.123| 0.111  | 0.052 | 0.64  | 0.527  | 0.2635|
| Experimental  | Time of day (minutes since midnight)   | 63  | 1    | 8    | 4.0  | 3.5    | 2.0   | 1.24  | 0.220  | 0.0853|
|               | Time since last meal (h)               | 52  | 0.061| 0.323| 0.123| 0.111  | 0.052 | 0.64  | 0.527  | 0.2635|
|               | Frame-wise displacement (mm)           | 63  | 0.061| 0.323| 0.123| 0.111  | 0.052 | 0.64  | 0.527  | 0.2635|
| Physiologic   | EGG frequency (Hz)                     | 63  | 0.041| 0.057| 0.049| 0.049  | 0.004 | 0.60  | 0.550  | 0.0000|
|               | EGG power (µV²)                        | 63  | 6    | 3056 | 281  | 107    | 479   | 0.18  | 0.858  | 0.0000|
|               | EGG cycle duration variability (s)     | 63  | 1    | 6.5  | 3    | 2.8    | 1.4   | 0.93  | 0.357  | 0.0605|
|               | LF-HRV power (ms²)                     | 52  | 0.5*—105| 4.5*—105| 9.5*—105| 0.7*—105| 0.7*—105| 3.76  | 0.0004 | 3099  |
|               | HF-HRV power (ms²)                     | 52  | 0.4*—105| 3.3*—105| 5.5*—105| 0.3*—105| 0.5*—105| 0.04  | 0.962  | 0.1463|
|               | LF/HF HRV ratio                        | 52  | 0.339| 8.15 | 2.51 | 1.91   | 1.97  | 1.87  | 0.067  | 0.3234|

For each variable tested, the table displays the descriptive statistics, t values, uncorrected p values, and Bayes factors (BFs), for which values smaller than 0.33 indicates substantial evidence for the null hypothesis. Gender effects were assessed by comparing males and females with an unpaired sample t-test, all other variables were assessed using robust linear regressions. Bold font indicates significant differences from the null distribution for p values and substantial evidence for the null hypothesis for bayes factors.
Possible functional roles of gastric-brain coupling

The precise function of the coupling between the BOLD signal and the gastric rhythm in this extended sensory and motor network remains highly speculative at this stage. Indeed, it takes a lot of information to define the function of a given brain region (Genon et al., 2018), let alone a whole network. Much remains to be studied, including notably task dependency, directionality of the interaction, as well as shared versus unique contribution of each sub-region of the gastric network. Still, the spatial layout of the gastric network and the involvement of all sensory and motor cortices is puzzling, and calls for further interpretation. In the following, we consider several nonexclusive working hypotheses regarding the potential functional consequences of gastric-brain coupling.

The gastric network could be functionally related to interoception and autonomic processes. Somatosensory, motor, premotor, cingulate motor and insular cortices, which belong to the gastric network, have established roles in interoception and autonomic processes (Critchley et al., 2004; Dum et al., 2016; Levinthal and Strick, 2020), and the strength of gastric-brain coupling is related to an index of cardiac activity. Given the physiological function of the stomach, a specific link with feeding behavior should be considered. A recent study found a negative correlation between weight loss and power at 0.05 Hz in gastric network regions across 90 individuals undergoing a weight reduction intervention (Levakov et al., 2021), suggesting a link between energy regulation and brain activity at gastric frequency. Note that the contribution of the cerebellum to the gastric network is compatible with a role in interoception and autonomic processes, since the cerebellum has been associated with activity from different bodily systems, such as cardiovascular (Reis and Golanov, 1997), respiratory (Xu and Frazier, 2000, p. 200), as well as in feeding control (Tataranni et al., 1999; Liu et al., 2000; Zhu and Wang, 2008). Still, the coupling between gastric rhythm and activity in early auditory and visual regions is not readily explained by a functional role of the gastric network limited to bodily regulations.
Garfinkel and Fries, 2005

spare most of the transmodal regions typically associated with arousal be concentrated in sensory and motor regions and the cardiac cycle (Garfinkel and Critchley, 2016), or to longer episodes of high or low arousal spanning several gastric cycles, remains to be determined. However, this view does not account for the specific layout of the gastric network. Indeed, why should gastric-related fluctuations of arousal be concentrated in sensory and motor regions and spare most of the transmodal regions typically associated with conceptual, abstract processing (Behrens et al., 2013)?

Finally, gastric-brain coupling might reflect an overall scaffolding mechanism contributing to the organization of large-scale neural activity, involved in the coordination between brain regions to bind information (Singer and Gray, 1995; Fries, 2005) expressed in different formats and coordinates (Azzalini et al., 2019). The sensory and motor regions of the gastric network contain topographical representations of the body surface, activated by touch, movement or visual perception of body parts (Penfield and Boldrey, 1937; Orlov et al., 2010; Amiez and Petrides, 2014), as well as of external space, which is represented in retinotopic coordinates in visual cortices and in tonotopic space in auditory cortices. The gastric rhythm, acting as a common input to all those regions, could facilitate the alignment and coordination of the different coordinate systems in which external information from the senses is expressed, in other words, act as a binding mechanism facilitating between-area communication (Azzalini et al., 2019), with the stomach delivering different time stamps to different regions in a mechanism reminiscent of traveling waves (Wang, 2010). Such a facilitation of interareal communication between sensory and motor regions would be an interesting complement to the known top-down control from cognitive areas to sensory regions.

In conclusion, the multiple cortical areas of the human brain have long been thought to be organized in an ascending hierarchy (Felleman and Van Essen, 1991) composed of relatively independent modules corresponding to each sensory modality converging onto higher-order, transmodal regions (Mesulam, 1998). While this view has been refined (Young, 1992; Markov et al., 2013), it is still much present in the narrative of large-scale brain organization, and fits with the classical parcellation in distinct resting-state networks for different modalities (Yeo et al., 2011). Our findings show that regions considered to be mostly independent are actually all tightly linked through delayed functional connectivity with the stomach. The monitoring of visceral inputs should thus be integrated into current views of the cortical hierarchy.

References

Addis DR, Wong AT, Schacter DL (2007) Remembering the past and imagining the future: common and distinct neural substrates during event construction and elaboration. Neuropsychologia 45:1363–1377.

Allen M, Poggiali D, Whitaker K, Marshall TR, Kiev RA (2019) Raincloud plots: a multi-platform tool for robust data visualization. Wellcome Open Res 4:63.

Amassian VE (1951) Cortical representation of visceral afferents. J Neurophysiol 14:433–444.

Amiez C, Petrides M (2014) Neuroimaging evidence of the anatomo-functional organization of the human cingulate motor areas. Cereb Cortex 24:563–578.

Andrews-Hanna JR, Smallwood J, Spreng RN (2014) The default network and self-generated thought: component processes, dynamic control, and clinical relevance. Ann NY Acad Sci 1316:29–52.

Ashburner J, Friston KJ (1999) Nonlinear spatial normalization using basis functions. Hum Brain Mapp 7:254–266.

Azzalini D, Rebollo I, Tallon-Baudry C (2019) Visceral signals shape brain dynamics and cognition. Trends Cogn Sci 23:488–509.

Barrett LF, Simmons WK (2015) Interceptive predictions in the brain. Nat Rev Neurosci 16:419–429.

Bastos AM, Schoffelen JM (2016) A tutorial review of functional connectivity analysis methods and their interpretational pitfalls. Front Syst Neurosci 9:175.

Behrens TEJ, Fox P, Laird A, Smith SM (2013) What is the most interesting part of the brain? Trends Cogn Sci 17:2–4.

Biswal B, Zerrin Yetkin F, Haughton VM, Hyde JS (1995) Functional connectivity in the motor cortex of resting human brain using echo-planar MRI. Magn Reson Med 34:537–541.

Cao J, Lu KH, Olesen ST, Phillips RJ, Jaffey D, Hendren CL, Powley TL, Liu Z (2019) Gastric stimulation drives fast BOLD responses of neural origin. Neuroimage 197:200–211.

Cappe C, Barone P (2005) Heteromodal connections supporting multisensory integration at low levels of cortical processing in the monkey. Eur J Neurosci 22:2886–2902.

Catani M (2017) A little man of some importance. Brain 140:3055–3061.

Cechetto DF, Saper CB (1987) Evidence for a viscerotopic sensory representation in the cortex and thalamus in the rat. J Comp Neurol 262:27–45.

Chang C, Metzger CD, Glover GH, Duyn JH, Heinez H-J, Walter M (2013) Association between heart rate variability and fluctuations in resting-state functional connectivity. Neuroimage 68:93–104.

Choe AS, Tang B, Smith KR, Honari H, Lindquist MA, Caffo BS, Pekar JJ (2020) Phase-locking of resting-state brain networks with the gastric basal electrical rhythm. bioRxiv. doi:10.1101/2020.06.32.085054.

Coen SJ, Hobson AR, Aziz Q (2012) Chapter 23—Processing of gastrointestinal sensory signals in the brain. In: Physiology of the gastrointestinal tract, Ed 5 (JohnsonLR, GhishanFK, KaunitzJD, MerchantJJ, SaidHM, and Wood JD, eds), pp 689–702. San Diego: Academic Press.

Critchley HD, Wiens S, Rotstein P, Öhman A, Dolan RJ (2004) Neural systems supporting interoceptive awareness. Nat Neurosci 7:189–195.

Cruz MT, Murphy EC, Sahibzada N, Verbalis JG, Gillis RA (2007) A reevaluation of the effects of stimulation of the dorsal motor nucleus of the vagus on gastric motility in the rat. Am J Physiol Regul Integr Comp Physiol 292:R291–R307.

de Munck JC, Gonçalves SL, Faes THc, Kuijer JPA, Pouwels PJW, Heeilha RM, Lopes da Silva FH (2008) A study of the brain’s resting-state based on spectral power, heart rate and fMRI. Neuroimage 42:112–121.

Delorme A, Makeig S (2004) EEGLAB: an open source toolbox for analysis of single-trial EEG dynamics including independent component analysis. J Neurosci Methods 134:9–21.

Desikan RS, Ségonne F, Fischl B, Quinn BT, Dickerson BC, Blacker D, Buckner RL, Dale AM, Maguire RP, Hyman BT, Albert MS, Killiany RJ (2006) An automated labeling system for subdividing the human cerebral cortex on MRI scans into gyral based regions of interest. Neuroimage 31:968–980.

Diedrichsen J, Balsters JH, Flavel J, Cussens E, Ramnani N (2009) A probabilistic MR atlas of the human cerebellum. Neuroimage 46:39–46.

Domschke K, Stevens S, Pfeiderer B, Gerlach AL (2010) Interoceptive sensitivity in anxiety and anxiety disorders: an overview and integration of neurobiological findings. Clin Psychol Rev 30:1–11.

Dum RP, Levinthal DJ, Strick PL (2009) The spinothalamic system targets motor and sensory areas in the cerebral cortex of monkeys. J Neurosci 29:14223–14235.

Dum RP, Levinthal DJ, Strick PL (2016) Motor, cognitive, and affective areas of the cerebral cortex influence the adrenal medulla. Proc Natl Acad Sci USA 113:9922–9927.

Elam M, Thorén P, Svensson TH (1986) Locus coeruleus neurons and sympathetic nerves activation by visceral afferents. Brain Res 375:117–125.

Eriq A, Van Horn SC, Sherman SM (1997) Relative numbers of cortical and brainstem inputs to the lateral geniculate nucleus. Proc Natl Acad Sci USA 94:1517–1520.

Felleman DJ, Van Essen DC (1991) Distributed hierarchical processing in the primate cerebral cortex. Cereb Cortex 1:1–47.

Felleman D, Van Essen DC (2005) Anatomy of Brain-Stomach Synchonosity.
Tataranni PA, Gautier J-F, Chen K, Uecker A, Bandy D, Salbe AD, Pratley RE, Lawson M, Reiman EM, Ravussin E (1999) Neuroanatomical correlates of hunger and satiation in humans using positron emission tomography. Proc Natl Acad Sci USA 96:4569–4574.

Tort ABL, Branka i/C20ck J, Draguhn A (2018) Respiration-entrained brain rhythms are global but often overlooked. Trends Neurosci 41:186–197.

Travagli RA, Hermann GE, Browning KN, Rogers RC (2006) Brainstem circuits regulating gastric function. Annu Rev Physiol 68:279–305.

Uhlrich DJ, Cucchiaro JB, Sherman SM (1988) The projection of individual axons from the parabrachial region of the brain stem to the dorsal lateral geniculate nucleus in the cat. J Neurosci 8:4565–4575.

Van Oudenhove J, Vandenberghe J, Dupont P, Geeraerts B, Vos R, Bormans G, Van Laere K, Fischler B, Demyttenaere K, Janssens J, Tack J (2009) Cortical deactivations during gastric fundus distension in health: visceral pain-specific response or attenuation of ‘default mode’ brain function? A H 2 15 O-PET study. Neurogastroenterol Motil 21:259–271.

Wagenmakers EJ (2007) A practical solution to the pervasive problems of p values. Psychon Bull Rev 14:779–804.

Wang XJ (2010) Neurophysiological and computational principles of cortical rhythms in cognition. Physiol Rev 90:1195–1268.

Wang Z, Bradesi S, Maarek JMI, Lee K, Winchester WJ, Mayer EA, Holschneider DP (2008) Regional brain activation in conscious, nonrestrained rats in response to noxious visceral stimulation. Pain 138:233–243.

Wetzels R, Wagenmakers EJ (2012) A default Bayesian hypothesis test for correlations and partial correlations. Psychon Bull Rev 19:1057–1064.

Wolpert N, Rebollo I, Tallon-Baudry C (2020) Electrogastrography for psychophysiological research: practical considerations, analysis pipeline, and normative data in a large sample. Psychophysiology 57:e13599.

Wu J, Ngo GH, Greve D, Li J, He T, Fischl B, Eickhoff SB, Yeo BTT (2018) Accurate nonlinear mapping between MNI volumetric and FreeSurfer surface coordinate systems. Hum Brain Mapp 39:3793–3808.

Xu F, Frazier DT (2000) Modulation of respiratory motor output by cerebellar deep nuclei in the rat. J Appl Physiol (1985) 89:996–1004.

Yeo BTT, Krienen FM, Sepulcre J, Sabuncu MR, Lashkari D, Hollinshead M, Roffman JL, Smoller JW, Zöllei L, Polimeni JR, Fischl B, Liu H, Buckner RL (2011) The organization of the human cerebral cortex estimated by intrinsic functional connectivity. J Neurophysiol 106:1125–1165.

Zhu JN, Wang JJ (2008) The cerebellum in feeding control: possible function and mechanism. Cell Mol Neurobiol 28:469–478.